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Published With international search report. (54) TIME: PETITIES AND COMPOUNDS THAT BIND TO A THROMBOPOIETIN RECEPTOR

#### (57) Abstract

Receptor are peptide and peptide mimeties that blad to and activate the thrombopoietin receptor. Such peptides and peptide mimeties are useful in methods for creating hematological disorders and particularly, thrombocytropenia realting from chemotherapy, addision therapy, or bone marrow transfusions as well as in diagnostic methods employing lateled peptides and peptide mimeties.

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#### PEPTIDES AND CCHPOUNDS THAT BIND TO A THRCHBOPOIETIN RECEPTOR

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# CROSS-REFERENCE TO RELATED CASES

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This application is a continuation-in-part of U.S. Patent Application Serial No. 08/485,301, filed June 7, 1995, and U.S. Patent Application Serial No. 08/478,128, filed June 7, 1995, each of which are herein incorporated by reference in their entirety for all purposes.

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## BACKGROUND OF THE INVENTION

The present invention provides peptides and compounds that bind to and activate the thrombopoietin receptor (c-mpl or TPO-R) or otherwise act as a TPO agonist. The invention has application in the fields of biochemistry and medicinal chemistry and particularly provides TPO agonists for use in the treatment of human disease.

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Megakaryocytes are bone marrow-derived cells, which are responsible for producing circulating blood platelets. Although comprising <0.25% of the bone marrow cells in most species, they have >10 times the volume of typical marrow cells. See Kuter et. al. Proc. Natl. Acad. Sci. USA 9::11104-11108 (1994). Megakaryocytes undargo a process known as endomitosis whereby they replicate their nuclei but fail to undergo cell division and thereby give rise to polyploid cells. In response to a decreased platelet count, the endomitotic rate increases, higher ploidy megakaryocytes are formed, and the number of megakaryocytes may increase up to 3-fold. See Harker 1. Clah. Invest. 47:458-465 (1968). In contrast, in response to an elevated platelet count, the

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endomitotic rate decreases, lower ploidy megakaryocytes are formed, and the number of megakaryocytes may decrease by 50%.

The exact physiological feedback mechanism by which e.g., Metcalf <u>Nature</u> 369:519-520 (1994). TPO has been shown platelets of recipient animals. Specifically, TPO is thought polyploidy, in megakaryocytes; (3) it increases megakaryocyte this feedback loop is now thought to be thrombopoletin (TPO) the mass of circulating platelets regulates the endomitotic More specifically, TPO has been shown to be the main humoral The circulating thrombopoietic factor involved in mediating rate and number of bone marrow megakaryocytes is not known. 'n. regulator in situations involving thrombocytopenia. See, in several studies to increase platelet counts, increase produces increases in megakaryocyte size and number; (2) to affect megakaryocytopoiesis in several ways: (1) it acetylcholinesterase-positive cells, in the bone marrow. platelet size, and increase isotope incorporation into megakaryocytes; and (5) it produces an increase in the endomitosis; (4) it produces increased maturation of produces an increase in DNA content, in the form of percentage of precursor cells, in the form of small

Because platelets (thrombocytes) are necessary for blood clotting and when their numbers are very low a patient is at serious risk of death from catastrophic hemorrhage, TPO has potential useful application in both the diagnosis and the treatment of various hematological disorders, for example, diseases primarily due to platelet defects. Ongoing clinical trials with TPO have indicated that TPO can be administered a basis for the projection of efficacy of TPO therapy in the treatment of thrombocytopenia, and particularly thrombocytopenia resulting from chemotherapy, radiation therapy, or bone marrow transplantation as treatment for cancer or lymphoma. See, e.g., McDonald (1992) Am. J. Ped. Hematology/Qncology 14:8-21 (1992).

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The gene encoding TPO has been cloned and characterized. See Kuter et al. <u>Proc. Natl. Acad. Sci. USA</u> 91:11104-11108 (1994); Barley et al. <u>Cell</u> 77:1117-1124

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(1994); Kaushanaky et al. Nature 369:568-571 (1994); Wendling et al. Nature 369:571-574 (1994); and Sauvage et al. Nature 369:533-538 (1994). Thrombopoietin is a glycoprotein with at least two forms, with apparent molecular masses of 25 kDa and 31 kDa, with a common N-terminal amino acid sequence. See, Bartley et al. Call 77:1117-1124 (1994). Thrombopoietin appears to have two distinct regions separated by a potential Arg-Arg cleavage site. The amino-terminal region is highly conserved in man and mouse, and has some homology with erythropoietin and interferon-a and interferon-b. The carboxy-terminal region shows wide species divergence.

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TPO-R is a member of the haematopoietin growth factor receptor The DNA sequences and encoded peptide sequences for residues in the N-terminal portion and a WSXWS motif close to USA 87:6934-6938 (1990). Evidence that this receptor plays a its expression is restricted to spleen, bone marrow, or fetal Vigon ec al. <u>Proc. Natl. Acad. Sci. USA</u> 89:5640-5644 (1992). megakaryocyte colonies without affecting erythroid or myeloid family, a family characterized by a common structural design (See Methia et al. <u>Blood</u> 82:1395-1401 (1993)). Furthermore, antisense to mpl RNA significantly inhibits the appearance of human TPO-R (also known as c-mpl) have been described. See See Bazan <u>Proc. Natl. Acad. Sci.</u> functional role in hematopoiesis includes observations that liver in mice (see Souyri et al. <u>Cell</u> 63:1137-1147 (1990)) and to megakaryocytes, platelets, and CD34 tells in humans colony formation. Some workers postulate that the receptor functions as a homodimer, similar to the situation with the of the extracellular domain, including four conserved C exposure of CD34 cells to synthetic oligonucleotides receptors for G-CSF and erythropoietin. the transmembrane region.

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The availability of cloned genes for TPO-R facilitates the search for agonists of this important receptor. The availability of the recombinant receptor protein allows the study of receptor-ligand interaction in a variety of random and semi-random peptide diversity generation systems. These systems include the "peptides on plasmids" system described in U.S. Patent Nos. 5,270,170 and 5,338,665;

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and in Cwirla et al., Proc. Natl. Acad. Sci. USA 87:6378-6382 Patent Application Serial No. 07/541,108, filed June 20, 1990, September 16, 1992, and 07/762,522, filed September 18, 1991; Fodor <u>Ann, Rep, Med. Chem.</u> 26:271-180 (1991); and U.S. Patent Application Serial No. 07/718,577, filed June 20, 1991, U.S. each of the foregoing patent applications and publications is Application Serial No. 08/300,262, filed September 2, 1994, 1990; Fodor et al. <u>Science</u> 251:767-773 (2/1991); Dower and Patent Application Serial No. 08/144,775, filed October 29, Publication No. 90/15070, published December 13, 1990; U.S. Patent Application Serial No. 07/624,120, filed December 6, Application Serial No. 07/805,727, filed December 6, 1991; which is a continuation-in-part application based on U.S. system described in U.S. Patent No. 5,143,854; PCT Patent 1993 and PCT WO 95/11992; the "encoded synthetic library" and the "very large scale immobilized polymer synthesis" the "peptides on phage" system described in U.S. Patent system described in U.S. Patent Application Serial Nos. (1990); the "polysome" system described in U.S. Patent 08/146,886, filed November 12, 1993, 07/946,239, filed incorporated herein by reference.

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The slow recovery of platelet levels in patients suffering from thrombocytopenia is a serious problem, and has lent urgency to the search for a blood growth factor agonist able to accelerate platelet regeneration. The present invention provides such an agonist.

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#### SUMMARY OF THE INVENTION

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This invention is directed, in part, to the novel and unexpected discovery that defined low molecular weight peptides and peptide mimetics have strong binding properties to the TPO-R and can activate the TPO-R. Accordingly, such peptides and peptide mimetics are useful for therapeutic purposes in treating conditions mediated by TPO (e.g., thrombocytopenia resulting from chemotherapy, radiation therapy, or bone marrow transfusions) as well as for diagnostic purposes in studying the mechanism of hematopoiesis

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and for the in vitro expansion of megakaroycytes and committed progenitor cells.

therapeutic and/or diagnostic purposes have an  ${\sf IC_{S0}}$  of about 2 purposes, the peptides and peptidomimetics preferably have an forth in Example 3 below wherein a lower  $\mathsf{IC}_{\mathsf{S0}}$  correlates to a than 500 nM. In a preferred embodiment, the molecular weight of the peptide or peptide mimetic is from about 250 to about mM or less, as determined by the binding affinity assay set  ${\sf IC}_{\sf SO}$  of no more than about 100  $\mu m$ , more preferably, no more stronger binding affinity to TPO-R. For pharmaceutical Peptides and peptide mimetics suitable for 8000 daltons.

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without such a label serve as intermediates in the preparation When used for diagnostic purposes, the peptides and peptide mimetics preferably are labeled with a detectable label and, accordingly, the peptides and peptide mimetics of labeled peptides and peptide mimetics.

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Weight and binding affinity for TPO-R comprise 9 or more amino Peptides meeting the defined criteria for molecular mimetics include peptides having one or more of the following synthetic (non-naturally occurring) amino acids. Pepcide acids wherein the amino acids are naturally occurring or modifications:

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linkages (bonds) have been replaced by a non-peptidyl linkage linkage; or an alkylated peptidyl linkage (-C(O) ${
m NR}^6$ - where  ${
m R}^6$ peptides wherein one or more of the peptidyl (-C(O) NR-) linkage; a urea [-NHC(O)NH-] linkage; a -CH2-secondary amine phosphonate linkage; a - $CH_2$ -sulfonamide [- $CH_2$ - $S(0)_2NR$ -] such as a -CH<sub>2</sub>-carbamate linkage (-CH<sub>2</sub>-OC(0)NR-); a is lower alkyl];

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Denzyloxycarbonyl-NH- group having from 1 to 3 substituents on peptides wherein the N-terminus is derivatized to a -NRR $^{
m l}$ hydrogen or lower alkyl with the proviso that R and R² are not -NRS(0) $_2$ R group; to a -NHC(0)NHR group where R and  $\mathbb{R}^1$  are group; to a -NRC(0)R group; to a -NRC(0)OR group; to a benzyloxycarbonyl-NH+ (CBZ-NH-) group; or to a both hydrogen; to a succinimide group; to a

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the phenyl ring selected from the group consisting of lower alkyl, lower alkoxy, chloro, and bromo;

peptides wherein the C terminus is derivatized to

 $-C(0)\,R^2$  where  $^2$  is selected from the group consisting of lower alkoxy, and -NR $^{\mathrm{J}}\mathrm{R}^{\mathrm{4}}$  where  $\mathrm{R}^{\mathrm{3}}$  and  $\mathrm{R}^{\mathrm{4}}$  are independently selected from the group consisting of hydrogen and lower alkyl.

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Accordingly, preferred peptides and peptide mimetics comprise a compound having:

(1) a molecular weight of less than about 5000

daltons, and

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(2) a binding affinity to TPO-R as expressed by an ICso of no more than about 100' μm,

wherein from zero to all of the -C(0)NH- linkages of the peptide have been replaced by a linkage selected from the group consisting of a

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-NHC(O)NH- linkage where R is hydrogen or lower alkyl and  $R^6$ -CH2OC(O)NR- linkage; a phosphonate linkage; a -CH2S(O)2NRlinkage; a -CH2NR- linkage; and a -C(O)NR6- linkage; and a is lower alkyl,

further wherein the N-terminus of said peptide or peptide group; a -NRC(O)R group; a -NRC(O)OR group; a -NRS(O)2R group; mimetic is selected from the group consisting of a -NRR1 a -NHC(O)NHR group; a succinimide group; a

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benzyloxycarbonyl-NH- group; and a benzyloxycarbonyl-NH- group chloro, and bromo, where R and R<sup>1</sup> are independently selected having from 1 to 3 substituents on the phenyl ring selected from the group consisting of lower alkyl, lower alkoxy, from the group consisting of hydrogen and lower alkyl,

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and still further wherein the C-terminus of said peptide and -NR<sup>3</sup>R<sup>4</sup> where R<sup>3</sup> and R<sup>4</sup> are independently selected from the nitrogen atom of the -NR3R4 group can optionally be the amine group of the N-terminus of the peptide so as to form a cyclic selected from the group consisting of hydroxy, lower alkoxy, group consisting of hydrogen and lower alkyl and where the or peptide mimetic has the formula -C(O)R2 where R2 is peptide,

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and physiologically acceptable salts thereof.

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In a related embodiment, the invention is directed to a labeled peptide or peptide mimetic comprising a peptide or peptide mimetic described as above having covalently attached thereto a label capable of detection.

In some embodiments of the invention, preferred peptides for use include peptides having a core structure comprising a sequence of amino acids:

genetically coded L-amino acids;  $X_{S}$  is A, D, E, G, K, M, Q, R, where X<sub>1</sub> is C, L, M, P, Q, V, X<sub>2</sub> is F, K, L, N, Q, R, S, T or S, T, V or Y; X<sub>6</sub> is C, F, G, L, M, S, V, W or Y; and X, is C, V; X3 is C, F, I, L, M, R, S, V or W; X4 is any of the 20 X1 X2 X3 X4 X5 X6 X7 G, I, K, L, M, N, R or V.

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In a preferred embodiment the core peptide comrpises a sequence of amino acids:

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is independently selected from any of the 20 genetically coded preferred embodiment,  $X_1$  is P;  $X_2$  is T;  $X_3$  is L;  $X_4$  is R;  $X_5$  is where  $X_1$  is L, M, P, Q, or V;  $X_2$  is F, R, S, or T;  $X_3$  is F, L, V, or W; X4 is A, K, L, M, R, S, V, or T; X5 is A, E, G, K, M, Q, R, S, or T; X, is C, I, K, L, M or V; and each Xg residue independently selected from any of the 20 genetically coded L-amino acids and their stereoisomeric D-amino acids. In a non-natural amino acids. Preferably, each  $X_{\rm B}$  residue is L-amino acids, their stereoisomeric D-amino acids; and XB G X1 X2 X3 X4X5 W X7 E or Q; and X, is I or L.

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More preferably, the core peptide comprises a

sequence of amino acids:

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is A, C, D, E, K, L, Q, R, S, T, or V. More preferably, X9 is where  $x_9$  is A, C, E, G, I, L , M, P, R, Q, S, T, or V; and  $x_8$ X9 X8 G X1 X2 X3 X4 X5 W X7 A or I; and Xg is D, E, or K. 30

Particularly preferred peptides include: G G C A D G LKSREHTS, SIE GPTLREWLTSRTPHS, LAIE N, G G C A D G P T L R E W I S F C G G K; T I K G P T L R Q W PTLREWISFCGG, GNADGPTLRQWLEGRRPK GPTLRQWLHGNGRDT; CADGPTLREWISFC; and I E G P T L R Q W L A A R A.

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peptides for use in this invention include peptides having a In further embodiments of the invention, preferred core structure comprising a sequence of amino acids:

C X2 X3 X4 X5 X6 X7

Y; and X, is C, G, I, K, L, M, N, R or V. In a more preferred V. Particularly preferred peptides include: G G C T L R E W where X<sub>2</sub> is K, L, N, Q, R, S, T or V; X<sub>3</sub> is C, F, I, L, M, R, In a further embodiment,  $X_2$  is S or T;  $X_3$  is L or R;  $X_4$  is R; S or V;  $X_4$  is any of the 20 genetically coded L-amino acids; Xs is A, D, E, G, S, V or Y; Xs is C, F, G, L, M, S, V, W or X<sub>5</sub> is D, E, or G; X<sub>6</sub> is F, L, or W; and X<sub>7</sub> is I, K, L, R, or embodiment, X4 is A, E, G, H, K, L, M, P, Q, R, S, T, or W. LHGG.FCGG.

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In a further embodiment, preferred peptides for use in this invention include peptides having a structure comprising a sequence of amino acids:

X<sub>8</sub> C X<sub>2</sub> X<sub>3</sub> X<sub>4</sub> X<sub>5</sub> X<sub>6</sub> X<sub>7</sub>

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acids; Xs is A, D, E, G, K, M, Q, R, S, T, V or Y; Xs is C, F, G, L, M, S, V, Wor Y; X, is C, G, I, K, L, M, N, R or V; and  $X_{8}$  is any of the 20 genetically coded L-amino acids. In some where X<sub>2</sub> is F, K, L, N, Q, R, S, T or V; X<sub>3</sub> is C, F, I, L, M, R, S, V or W; X4 is any of the 20 genetically coded L-amino embodiments, Xg is preferably G, S, Y, or R.

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particularly for treating hematological disorders, including chemotherapy, radiation therapy, or bone marrow transfusions The compounds described herein are useful for the administered, a therapeutically effective dose or amount of 18 prevention and treatment of diseases mediated by TPO, and Busceptible to treatment with a TPO agonist receives, or Thus, the present invention also provides a method for treating wherein a patient having a disorder that is but not limited to, thrombocytopenia resulting from compound of the present invention.

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compositions comprising one or more of the compounds described The invention also provides for pharmaceutical

including oral dosage forms, as well as inhalable powders and pharmaceutical compositions can be in a variety of forms These herein and a physiologically acceptable carrier. solutions and injectable and infusible solutions.

# BRIEF DESCRIPTION OF THE FIGURES

Figures 1A-B illustrates the results of a functional described in Example 2. Figure 1A is a graphical depiction of the results of the TPO-R transfected Ba/F3 cell proliferation assay in the presence of various peptides; the assay is assay for selected peptides of the invention:

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designating the results for G G C A D G P T L R E W ISFCGGK (biotin); Ġ

designating the results for G G C A D G P T L R E W ISFCGG; ×

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designating the results for LAIEGPTLROWL HGNGRDT;

designating the results for GNADGPTLROW EGRRPKN; and o

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designating the results for TIKGPTLROWLK SREHTS.

Figure 1B is a graphical depiction of the results with the same peptides and the parental cell line. Figure 2A-C show the results of peptide

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Figure 2A shows the results of the assay Figure 2C shows the results of the assay streptavidin (SA)) for both the transfected and parental cell lines. Figure 2B shows the results of the assay for the free biotinylated peptide (AF 12285) for both the transfected and for streptavidin alone for both the transfected and parental oligomerization using the TPO-R transfected Ba/F3 cell for the complexed biotinylated peptide (AF 12285 with proliferation assay. parental cell lines. cell lines.

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Figures 3A-G show the results of a series of control experiments showing the activity of TPO, the peptides of the present invention, EPO, and EPO-R binding peptides in a cell proliferation assay using either the TPO-R transfected Ba/F3

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proliferation assay using the EPO-dependent cell line. Figure 3F depicts the results for EPO in the cell proliferation assay the results for complexed biotinylated peptide (AF 12285 with line and its corresponding parental line. Figure 3C depicts EPO-dependent cell line. Figure 3A depicts the results for proliferation assay using the TPO-R transfected Ba/F3 cell transfected Ba/F3 cell line and its corresponding parental streptavidin (SA)) and a complexed form of a biotinylated using the EPO-dependent cell line. Figure 3G depicts the results for complexed biotinylated peptide (AF 12885 with line. Figure 3B depicts the results for EPO in the cell corresponding parental cell line are shown in Figure 3D. EPO-R binding peptide (AF 11505 with SA) in the TPO-R cell line and its corresponding parental line, or an TPO in the cell proliferation assay using the TPO-R transfected Ba/F3 cell line. The results for the Figure 3E depicts the results for TPO in the cell w

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EPO-R binding peptide (AF 11505 with SA) in the EPO-dependent streptavidin (SA)) and the complexed form of a biotinylated cell line. 15 20

intragenic region (M13 IG) to permit rescue of single-stranded the bla gene to permit selection on ampicillin, the M13 phage library plasmid includes the  $\it rrnB$  transcriptional terminator, DNA, a plasmid replication origin (ori), two  $laco_s$  sequences, and the arac gene to permit positive and negative regulation gene. Figure 4B shows the sequence of the cloning region at sites used during library construction. Figure 4C shows the peptides-on-plasmids libraries in vector pJS142. Figure 4A the 3' end of the lac I gene, including the Sfil and Eagl of the araB promoter driving expression of the lac fusion ligation of annealed library oligonucleotides, ON-829 and shows a restriction map and position of the genes. The Figures 4A-C illustrates the construction of ON-830, to Sfil sites of pJS142 to produce a library. spaces in the sequence indicate sites of ligation.

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Figures 5A-B illustrate cloning into the pELM3 and PELM15 MBP vectors. Figure 5A shows the sequence at the 3' and of the malE fusion gene, including the MBP coding

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sequence, the poly asparagine linker, the factor Xa protease cleavagge site, and the available cloning sites. The remaining portions of the vectors are derived from pMALC2 (PELM3) and pMALD2 (PELM15), available from New England Biolabs. Figure 5B shows the sequence of the vectors after transfer of the BspEII-ScaI library fragment into AgeI-ScaI digested pELM3/pELM15. The transferred sequence includes the sequence encoding the GGG peptide linker from the pJS142 library.

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Figure 6A depicts a restriction map and position of the genes for the construction of headpiece dimer libraries in vector pCMG14. The library plasmid includes: the *xrnB* transcriptional terminator, the *bla* gene to permit selection on ampicillin, the M13 phage intragenic region (M13 IG) to permit rescue of single-stranded DNA, a plasmid replication origin (ori), one laco<sub>g</sub> ssequence, and the araC gene to permit positive and negative regulation of the araB promoter driving expression of the headpiece dimer fusion gene. Figure 6B depicts the sequence of the cloning region at the 3' end of the headpiece dimer gene, including the Sfil and Eagl sites used during library construction. Figure 6C shows the ligation of annealed ON-1679, ON-829, and ON-830 to Sfil sites of pCMG14 to produce a library. Singles spaces in the sequence indicate sites of ligation.

Figures 7 to 9 show the results of further assays evaluating activity of the peptides and peptide mimETICS of the invention. In this assay mice are made thrombocytopenic with carboplatin. Figure 7 depicts typical results when balb/C mice are treated with carboplatin (125 mg/kg intraperitoneally) on Day 0. The dashed lines represent untreated animals from three experiments. The solid line represent carboplatin-treated groups in three experiments. The heavy solid lines represent historical data. Figure 8 depicts the effect of carboplatin titration on platelet counts in mice treated with the indicated amounts of carboplatin (in mg/kg, intraperitoneally (ip) on Day 0). Figure 9 depicts amelioration of carboplatin-induced thrombocytopenia on Day 10 by peptide AF12513 (513). Carboplatin (CBP; 50-125 mg/kg,

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intraperitoneally) was administered on Day 0. AF12513 (1 mg/kg, ip) was given on Days 1-9.

# DESCRIPTION OF SPECIFIC FABODIMENTS

# DEPINITIONS AND GENERAL PARAMETERS

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The following definitions are set forth to illustrate and define the meaning and scope of the various terms used to describe the invention herein.

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"Agonist" refers to a biologically active ligand which binds to its complementary biologically active receptor and activates the latter either to cause a biological response in the receptor or to enhance preexisting biological activity of the receptor.

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"Pharmaceutically acceptable salts" refer to the non-toxic alkali metal, alkaline earth metal, and ammonium salts commonly used in the pharmaceutical industry including the sodium, potassium, lithium, calcium, magnesium, barium, ammonium, and protamine zinc salts, which are prepared by methods well known in the art. The term also includes non-toxic acid addition salts, which are generally prepared by reacting the compounds of this invention with a suitable organic or inorganic acid. Representative salts include the hydrochloride, hydrobromide, sulfate, bisulfate, acetate, oxalate, valerate, oleate, laurate, borate, benzoate, lactate, phosphate, tosylate, citrate, maleate, fumarate, succinate, tattrate, napsylate, and the like.

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"Pharmaceutically acceptable acid addition salt" refers to those salts which retain the biological effectiveness and properties of the free bases and which are not biologically or otherwise undesirable, formed with inorganic acids such as hydrochloric acid, hydrobromic acid, sulfuric acid, phosphoric acid and the like, and organic acids such as acetic acid, propionic acid, glycolic acid, pyruvic acid, oxalic acid, malic acid, malonic acid, succinic acid, maleic acid, fumaric acid, tartaric acid, citric acid, benzoic acid, cinnamic acid, mandelic acid, citric acid, mandelic acid,

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menthanesulfonic acid, ethanesulfonic acid, p-toluenesulfonic acid, salicylic acid and the like. For a description of pharmaceutically acceptable acid addition salts as prodrugs, see Bundgaard, H., supra.

biological effectiveness and properties of the carboxylic acid e.g., March <u>Advanced Organic Chemistry</u>, 3rd Ed., John Wiley & "Pharmaceutically acceptable ester" refers to those described herein and at the same time are the pharmaceutically or alcohol and are not biologically or otherwise undesirable. Wiley & Sons, New York (1980)). The alcohol component of the or heteroaromatic alcohols. This invention also contemplates or can not contain branched carbons or (ii) a  $C_7$ - $C_{12}$  aromatic Elsevier Science Publishers, Amsterdam (1985). These esters that can or can not contain one or more double bonds and can esters which retain, upon hydrolysis of the ester bond, the are typically formed from the corresponding carboxylic acid ester will generally comprise (i) a  $C_2$ - $C_{12}$  aliphatic alcohol Sons, New York (1985) p. 1157 and references cited therein, For a description of pharmaceutically acceptable esters as and Mark et al. Encyclopedia of Chemical Technology, John the use of those compositions which are both esters as prodrugs, see Bundgaard, H., ed., <u>Dasion of Prodrugs</u>, accomplished via conventional synthetic techniques. Generally, ester formation can be acceptable acid addition salts thereof. and an alcohol.

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"Pharmaceutically acceptable amide" refers to those amides which retain, upon hydrolysis of the amide bond, the biological effectiveness and properties of the carboxylic acid or amine and are not biologically or otherwise undesirable. For a description of pharmaceutically acceptable amides as produces, see Bundgaard, H., ed., Design of Produces as produces, see Bundgaard, H., ed., Design of Produces as are typically formed from the corresponding carboxylic acid and an amine. Generally, amide formation can be accomplished via conventional synthetic techniques. (See, e.g., March Advanced Organic Chemistry, 1rd Ed., John Wiley & Sons, New York (1985) p. 1152 and Mark et al. Encyclopedia of Chemical Technology, John Wiley & Sons, New York (1985)). This

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invention also contemplates the use of those compositions which are both amides as described herein and at the same time are the pharmaceutically acceptable acid addition salts thereof.

"Pharmaceutically or therapeutically acceptable carrier" refers to a carrier medium which does not interfere with the effectiveness of the biological activity of the active ingredients and which is not toxic to the host or patient.

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"Stereoisomer" refers to a chemical compound having the same molecular weight, chemical composition, and constitution as another, but with the atoms grouped differently. That is, certain identical chemical moieties are at different orientations in space and, therefore, when pure, has the ability to rotate the plane of polarized light. However, some pure stereoisomers may have an optical rotation that is so slight that it is undetectable with present instrumentation. The compounds of the instant invention may have one or more asymmetrical carbon atoms and therefore include various stereoisomers. All stereoisomers are included within the scope of the invention.

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"Therapeutically- or pharmaceutically-effective amount" as applied to the compositions of the instant invention refers to the amount of composition sufficient to induce a desired biological result. That result can be alleviation of the signs, symptoms, or causes of a disease, or any other desired alteration of a biological system. In the present invention, the result will typically involve a decrease in the immunological and/or inflammatory responses to infection or tissue injury.

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Amino acid residues in peptides are abbreviated as follows: Phenylalanine is Phe or F; Leucine is Leu or L; Isoleucine is Ile or I; Methionine is Met or M; Valine is Val or V; Serine is Ser or S; Proline is Pro or P; Threonine is Thr or T; Alanine is Ala or A; Tyrosine is Tyr or Y; Histidine is His or H; Glutamine is Gln or Q; Asparagine is Asn or N; Lysine is Lys or K; Aspartic Acid is Asp or D; Glutamic Acid is Glu or E; Cysteine is Cys or C; Tryptophan is Trp or W;

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Thi is thienylalanine, OBn is O-benzyl, and hyp is

hydroxyproline.

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types of non-peptide compound are termed "peptide mimetics" or reference). Peptide mimetics that are structurally similar to naturally-occurring receptor-binding polypeptide, but have one "Peptidomimetics" (Fauchere, J. Adv. Drug Rgs. 15:29 (1986); analogs are also provided. Peptide analogs are commonly used Trends Pharm Sci. (1980) pp. 463-468 (general review); Hudson, -CH<sub>2</sub>SO-, by methods known in the art and further described in -CH<sub>2</sub>-CH<sub>2</sub>-, -CH=CH- (cis and trans), -COCH<sub>2</sub>-, -CH(OH)CH<sub>2</sub>-, and (-CH2-S); Hann J. Chem. Soc .Perkin Trans. J. 307-314 (1982) naturally-occurring amino acids, peptidomimetics or peptide Veber and Freidinger IINS p.392 (1985); and Evans et al. <u>J.</u> Med. Chem. 30:1229 (1987), which are incorporated herein by D. et al., Int J Pept Prot Res 14:177-185 (1979) (-CH2NH-, therapeutically useful peptides may be used to produce an equivalent or enhanced therapeutic or prophylactic effect, or more peptide linkages optionally replaced by a linkage the following references: Spatola, A.F. in Chemistry and in the pharmaceutical industry as non-peptide drugs with Generally, peptidomimetics are structurally similar to a Reptide Backbone Modifications (general review); Morley, selected from the group consisting of:  $^- ext{CH}_2 ext{NH}^-$ ,  $^- ext{CH}_2 ext{S}^-$ , Weinstein, eds., Marcel Dekker, New York, p. 267 (1983); letxahedron Lett 23:2533 (1982) (-COCH2-); Szelke et al. properties analogous to those of the template peptide. (-CH-CH-, cis and trans); Almquist et al. J. Med. Chem. Biochemistry of Amino Acids, Peptides, and Proteins, B. Spatola, A.F., <u>Vega Data</u> (March 1983), Vol. 1, Issue 3, In addition to peptides consisting only of paradigm polypeptide (i.e., a polypeptide that has a CH<sub>2</sub>CH<sub>2</sub>-); Spatola et al. <u>Life Sci</u> 38:1243-1249 (1986) 13:1392-1398 (1980) (-COCH<sub>2</sub>-); Jennings-White et al. biological or pharmacological activity), such as

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A particularly preferred non-peptide linkage is -CH2NH-. Such (-CH(OH)CH2-); Holladay et al. Tetrahedron Lett 24:4401-4404 attachment of one or more labels, directly or through a spacer  $(-CH_2-S-)$ ; each of which is incorporated herein by reference. (e.g., an amide group), to non-interfering position(s) on the non-interfering positions generally are positions that do not Derivitization (e.g., labeling) of peptidomimetics should not receptor with high affinity and possess detectable biological efficacy, etc.), altered specificity (e.g., a broad-spectrum of biological activities), reduced antigenicity, and others. activity (i.e., are agonistic or antagonistic to one or more economical production, greater chemical stability, enhanced pharmacological properties (half-life, absorption, potency, Generally, (1983) (-C(OH)CH2-); and Hruby Life Sci 31:189-199 (1982) structure-activity data and/or molecular modeling. Such peptidomimetics of receptor-binding peptides bind to the Peptidomimetic binds to produce the therapeutic effect. form direct contacts with the macromolecules(s) (e.g., substantially interfere with the desired biological or peptide mimetics may have significant advantages over polypeptide embodiments, including, for example: more Labeling of peptidomimetics usually involves covalent European Appln. EP 45665 CA (1982): 97:39405 (1982) immunoglobulin superfamily molecules) to which the Peptidomimetic that are predicted by quantitative pharmacological activity of the peptidomimetic. receptor-mediated phenotypic changes).

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consensus sequence variation may be generated by methods known (e.g., D-lysine in place of L-lysine) may be used to generate of a consensus sequence with a D-amino acid of the same type Systematic substitution of one or more amino acids comprising a consensus sequence or a substantially identical intramolecular disulfide bridges which cyclize the peptide (1992), incorporated herein by reference); for example, by more stable peptides. In addition, constrained peptides in the art (Rizo and Gierasch Ann. Rev. Biochem, 61:387 adding internal cysteine residues capable of forming

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atoms, 3) a heterocycle of from 3 to 7 carbon atoms and 1 to 2 carbon atoms optionally having from 1 to 3 substituents on the refer to amino acids which do not naturally occur in vivo but structures described herein. Preferred synthetic amino acids are the D-a-amino acids of naturally occurring L-a-amino acid  $R^2$  is selected from the group consisting of hydrogen, hydroxy, hydroxy, (b) amino, (c) cycloalkyl and cycloalkenyl of from 3 nucleus selected from the group consisting of hydroxyl, lower consisting of oxygen, sulfur, and nitrogen, (f) -C(0)R2 where represented by the formula H2NCHR5COOH where R5 is 1) a lower sulfur, and nitrogen, 4) an aromatic residue of from 6 to 10 and lower alkyl, (g)  $-S(0)_nR^6$  where n is an integer from 1 to 2 and R<sup>6</sup> is lower alkyl and with the proviso that R<sup>5</sup> does not independently selected from the group consisting of hydrogen hydroxyl, lower alkoxy, amino, and carboxyl, 5) -alkylene-Y alkoxy, amino and carboxyl, (e) heterocyclic of from 3 to 7 carbon atoms and 1 to 2 heteroatoms selected from the group optionally having from 1 to 3 substituents on the aromatic which, nevertheless, can be incorporated into the peptide as well as non-naturally occurring D- and L-a-amino acids heteroatoms selected from the group consisting of oxygen, where alkylene is an alkylene group of from 1 to 7 carbon Synthetic or non-naturally occuring amino acids alkyl group, 2) a cycloalkyl group of from 3 to 7 carbon lower alkyl, lower alkoxy, and -NR<sup>3</sup>R<sup>4</sup> where R<sup>3</sup> and R<sup>4</sup> are atoms and Y is selected from the group consisting of (a) to 7 carbon atoms, (d) aryl of from 6 to 10 carbon atoms define a side chain of a naturally occurring amino acid. aromatic nucleus selected from the group consisting of

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Other preferred synthetic amino acids include amino acids wherein the amino group is separated from the carboxyl group by more than one carbon atom such as b-alanine, g-aminobutyric acid, and the like.

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L-2-naphthylalanine, L-cyclohexylalanine, L-2-amino isobutyric include, by way of example, the D-amino acids of naturally acid, the sulfoxide and sulfone derivatives of methionine Particularly preferred synthetic amino acids occurring L-amino acids, L-1-napthyl-alanine,

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methionine (1.e., HOOC-(H2NCH)CH2CH2-OR6 where R6 is as defined (i.e., HOOC- $(H_2NCH) CH_2CH_2-S(O)_nR^6$ ) where n and  $R_6$  are as defined above as well as the lower alkoxy derivative of above).

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mimetics in vivo in the patient to whom the peptide or peptide mimetic has been administered. Suitable detectable labels are radioisotopes, fluorescent labels (e.g., fluorescein), and the critical and is selected relative to the amount of label to be employed as well as the toxicity of the label at the amount of Detectable label" refers to materials, which when covalently attached to the peptides and peptide mimetics of this invention, permit detection of the peptide and peptide label employed. Selection of the label relative to such like. The particular detectable label employed is not well known in the art and include, by way of example, factors is well within the skill of the art.

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chemistry. Likewise, <sup>32</sup>P can be incorporated onto the peptide or peptide mimetic as a phosphate moiety through, for example, be achieved by incorporating the amino acid tyrosine into the peptide or peptide mimetic and then iodating the peptide. If Covalent attachment of the detectable label to the attachment of  $^{125}\mathrm{I}$  to the peptide or the peptide mimetic can peptide or peptide mimetic is accomplished by conventional radioisotope is employed as the detectable label, covalent methods well known in the art. For example, when the  $^{125}\mathrm{I}$ tyrosine is not present in the peptide or peptide mimetic, peptide or peptide mimetic can be achieved by well known a hydroxyl group on the peptide or peptide mimetic using incorporation of tyrosine to the N or C terminus of the conventional chemistry.

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#### II. OVERVIEW

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agonist. These compounds include "lead" peptide compounds and similar molecular structure or shape as the lead compounds but "derivative" compounds constructed so as to have the same or The present invention provides compounds that bind to and activate the TPO-R or otherwise behave as a TPO

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that differ from the lead compounds either with respect to susceptibility to hydrolysis or proteolysis and/or with respect to other biological properties, such as increased affinity for the receptor. The present invention also provides compositions comprising an effective amount of a TPO agonist, and more particularly a compound, that is useful for treating hematological disorders, and particularly, thrombocytopenia associated with chemotherapy, radiation therapy, or bone marrow transfusions.

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# III. <u>IDENTIFICATION OF TPO-AGONISTS</u>

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Peptides having a binding affinity to TPO-R can be readily identified by random peptide diversity generating systems coupled with an affinity enrichment process.

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systems include the "peptides on plasmids" system described in immobilized polymer synthesis" system described in U.S. Patent Application Serial No. 08/144,775, filed October 29, 1993 and PCT WO 95/11992; the "encoded synthetic library (ESL)" system phage" system described in U.S. Patent Application Serial No. No. 5,143,854; PCT Patent Publication No. 90/15070, published application of U.S. Patent Application Serial No. 07/946,239, application of U.S. Patent Application Serial No. 07/762,522, 07/541,108, filed June 20, 1990, and in Cwirla et al., <u>Proc.</u> Specifically, random peptide diversity generating described in U.S. Patent Application Serial No. 08/146,886, U.S. Patent Nos. 5,270,170 and 5,338,665; the "peptides on 07/718,577, filed June 20, 1991 which is a continuation in filed September 16, 1992, which is a continuation in part 07/624,120, filed December 6, 1990; Fodor et al. <u>Srience</u> Natl. Agad. Sci. USA 87:6378-6382 (1980); the "polysome filed November 12, 1993 which is a continuation in part system" described in U.S. Patent Application Serial No. part application of U.S. Patent Application Serial No. continuation-in-part application based on U.S. Patent December 13, 1990; U.S. Patent Application Serial No. filed September 18, 1991; and the "very large scale 08/300,262, filed September 2, 1994, which is a

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251:767-773 (2/1991); Dower and Fodor <u>Ann. Rep. Med. Chem.</u> 26:271-180 (1991); and U.S. Patent Application Serial No. 805,727, filed December 6, 1991.

Using the procedures described above, random Peptides were generally designed to have a defined number of amino acid residues in length (e.g., 12). To generate the collection of oligonucleotides encoding the random peptides, the codon motif (NNK)x, where N is nucleotide A, C, G, or T (equimolar; depending on the methodology employed, other nucleotides can be employed), K is G or T (equimolar), and x is an integer corresponding to the number of amino acids in the peptide (e.g., 12) was used to specify any one of the 32 possible codons resulting from the NNX motif: 1 for each of 12 amino acids, 2 for each of 5 amino acids, 3 for each of 3 amino acids, and only one of the three stop codons. Thus, the NNX motif encodes all of the amino acids, encodes only one stop codon, and reduces codon bias.

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In the systems employed, the random peptides were presented either on the surface of a phage particle, as part of a fusion protein comprising either the pIII or the pVIII coat protein of a phage fd derivative (peptides on phage) or as a fusion protein with the LacI peptide fusion protein bound to a plasmid (peptides on plasmids).

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The phage or plasmids, including the DNA encoding the peptides, were identified and isolated by an affinity enrichment process using immobilized TPO-R. The affinity enrichment process, sometimes called 'panning," involves multiple rounds of incubating the phage, plasmids, or polysomes with the immobilized receptor, collecting the phage, plasmids, or polysomes that bind to the receptor (along with the accompanying DNA or mRNA), and producing more of the phage or plasmids (along with the accompanying LacI-peptide fusion protein) collected. The extracellular domain (ECD) of the TPO-R typically was used during panning.

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After several rounds of affinity enrichment, the phage or plasmids and accompanying peptides were examined by ELISA to determine if the peptides bind specifically to TPO-R. This assay was carried out similarly to the procedures used in

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A similar ELISA procedure for use in the pepides on alkaline phosphatase in each well was determined by standard the affinity enrichment process, except that after removing unbound phage, the wells were typically treated with rabbit (AP) -conjugated goat anti-rabbit antibody. The amount of anti-phage antibody, then with alkaline phosphatase plasmids system is described in detail below.

radiolabelled monovalent receptor. This probe can be produced receptor), one can determine whether the fusion proteins bind TPO-R specific phage clones. The receptor is then labeled to host cells, typically CHO cells. Following PI-PLC harvest of high specific activity with  $^{33}\mathrm{P}$  for use as a monovalent probe to the receptor specifically. The phage pools found to bind to TPO-R were screened in a colony lift probing format using the receptors, the receptor was tested for binding to TPO or using protein kinase A to phosphorylate a kemptide sequence "engineered" form of the TPO receptor is then expressed in By comparing test wells with control wells (no fused to the C-terminus of the soluble receptor. The to identify high affinity ligands using colony lifts.

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were then synthesized as the free peptide (e.g., no phage) and peptide portion that are preferred compounds of the invention. Peptides found to bind specifically to the receptor proteins for which the binding to the receptor was blocked by tested in a blocking assay. The blocking assay was carried TPO or the reference peptide contain peptides in the random out in similar fashion to the ELISA , except that TPO or a reference peptide was added to the wells before the fusion receptor; and (2) no TPO or reference peptide). Fusion protein (the control wells were of two types: (1) no

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produced in recombinant host cells. One useful form of TPO-R is constructed by expressing the protein as a soluble protein in baculovirus transformed host cells using standard methods; another useful form is constructed with a signal peptide for protein secretion and for glycophospholipid membrane anchor TPO-R, as well as its extracellular domain, were attachment. This form of anchor attachment is called

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"PIG-tailing". See Caras and Wendell <u>Science</u> 243:1196-1198 (1989) and Lin et al. Science 249:677-679 (1990).

receptor from the surface of the cells expressing the receptor immobilizing antibody to block unbound sites that remain after expression of receptor with a cell sorter) with phospholipase signal protein for membrane attachment and can be immobilized protein can be immobilized by coating the wells of microtiter immobilization reaction in varying concentrations of receptor addition, one should ensure that the immobilizing antibody is during the affinity enrichment process. Otherwise, unblocked completely blocked (with TPO or some other blocking compound) blocking non-specific binding with bovine serum albumin (BSA) in PBS, and then binding cleaved recombinant receptor to the enrichment procedure. One can use peptides that bind to the antibody. See U.S. Patent Application Serial No. 07/947,339, C. The cleaved receptor still comprises a carboxy terminal plates with an anti-HPAP tail antibody (Ab 179 or MAb 179), because different preparations of recombinant protein often Using the PIG-tailing system, one can cleave the filed September 18, 1992, incorporated herein by reference. sequence of amino acids, called the "HPAP tail", from the to determine the optimum amount for a given preparation, receptor immobilization to avoid this problem or one can without further purification. The recombinant receptor antibody. Using this procedure, one should perform the simply immobilize the receptor directly to the wells of contain different amounts of the desired protein. In antibody can bind undesired phage during the affinity microtiter plates, without the aid of an immobilizing (e.g., transformed CHO cells selected for high level 'n ដ

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that can bind to the immobilized receptor. At higher receptor treated with 0.25 to 0.5 mg of receptor), multivalent binding recognize that the density of the immobilized receptor is an When using random peptide generation systems that allow for multivalent ligand-receptor interaction, one must important factor in determining the affinity of the ligands densities (e.g., each anti-receptor antibody-coated well is more likely to occur than at lower receptor densities

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monovalent receptor probe frequently is used. This probe can be produced using protein kinase A to phosphorylate a kemptide host cells, typically CHO cells. Following PI-PLC harvest of TPO-R specific phage clones. The receptor is then labeled to high specific activity with <sup>13</sup>P for use as a monovalent probe the receptors, the receptor was tested for binding to TPO or To discriminate among higher affinity peptides, a "engineered" form of the TPO receptor is then expressed in sequence fused to the C-terminus of the soluble receptor. to identify high affinity ligands using colony lifts.

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binds to TPO-R. The phage DNAs are sequenced to determine the PVIII-based peptides on phage system, a random library can be domain of the receptor and then making other peptides which resemble the lead peptides. Specifically, using a pIII or screened to discover a phage that presents a peptide that identifying lead peptides which bind to the extracellular sequences of the peptides displayed on the surface of the identification of peptides which bind TPO-R involve first Preferred screening methods to facilitate phages.

Clones capable of specific binding to the TPO-R were construction of other peptide libraries designed to contain a peptides. These libraries can be synthesized so as to favor peptide in only a few residues. This approach involves the identified from a random linear 10-mer pVIII library and a high frequency of derivatives of the initially identified synthesis of an oligonucleotide with the binding peptide the production of peptides that differ from the binding sequences of these peptides serve as the basis for the random cyclic 10-mer and 12-mer pVIII libraries. The coding sequence, except that rather than using pure

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mixture for this purpose) so as to generate derivatives of the triphosphates (i.e., 55% of the "correct" nucleotide, and 15% preparations of each of the four nucleoside triphosphates in each of the other three nucleotides is one preferred mixture for this purpose and 70% of the "correct" nucleotide and 10% of each of the other three nucleotides is another preferred the synthesis, one uses mixtures of the four nucleoside binding peptide coding sequence.

clones which enclode the sequence XXXXX (C, S, P, or R) TLREWL A variety of strategies were used to derivatize the XXXXXXX. An additional extended/mutagenized library, XXXX (C, clones which enclode the sequence XXXX (C, S, P, or R) TLREWL and extended on each terminus with random residues to produce S, P, or R) TIREWL XXXXXX (C or S), was constructed using the These included a pVIII phagemid mutagenesis library based on XXXXXX (C or S). A similar extended/mutagenized library was constructed using the peptides-on-plasmids system to produce lead peptides by making "mutagenesis on a theme" libraries,. with peptide elution and probed with radiolabeled monovalent the consensus sequence mutagenized at 70:10:10:10 frequency polysome display system. All three libraries were screened receptor.

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for peptide screening and mutagenesis studies and is described The plasmids thus isolated can then be reintroduced The "peptides on plasmids" techniques was also used incorporated herein by reference for all purposes. According to this approach, random peptides are fused at the C-terminus of LacI through expression from a plasmid vector carrying the screened by affinity purification (panning) on an immobilized encoding DNA occurs via the laco sequences on the plasmid, forming a stable peptide-LacI-plasmid complex that can be in greater detail in U.S. Patent no. 5,338,665, which is population for additional rounds of screening, or for the fusion gene. Linkage of the LacI-peptide fusion to its into E. coli by electroporation to amplify the selected examination of individual clones. receptor.

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mutagenesis studies were performed using a modified C-terminal In addition, random peptide screening and

expression as a C-terminal fusion protein. Crude cell lysates screened and the resulting DNA inserts were cloned as a pool from randomly picked individual MBP fusion clones were then assayed for TPO-R binding in an ELISA format, as discussed into a maltose binding protein (MBP) vector allowing their Lac-I display system in which display valency was reduced The libraries were ("headpiece dimer" display system).

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70:10:10:10 mutagenesis at positions 1 and 2 and K (G or T) at using the polysome display system, as described in co-pending A mutagenesis library was constructed based on the position 3 of the codon. The library was panned for 5 rounds 08/144,775, filed October 29, 1993 and PCT WO 95/11992, each and their binding affinities were determined by an MBP ELISA. sequence X X X X (C,P,R,or S) t l r e f l X X X X X (C or S), in which X represents a random NNK codon, and the lower The sequences were subcloned into an MBP vector against TPO receptor which had been immobilzed on magnetic application U.S. Patent Application Serial No. 08/300,262, Peptide mutagenesis studies were also conducted After the fifth round, the PCR amplified pool was filed September 2, 1994, which is a continuation-in-part application based on U.S. Patent Application Serial No. of which is incorporated herein by references for all cloned into pAFF6 and the ELISA positive clones were case letters represent amino acid codons containing sequenced. peads.

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magnetic beads (available from Dynal Corporation) as described in 0.5 M borate buffer (pH 9.5) overnight at room temperature by the manufacturer. The beads were incubated with antibody The beads were washed and combined with TPO-R containing the incubated for 1 hour at 4°C, and the beads were washed again To imobilize the TPO-R for polysome screening, Ab "HPAP" tail. The antibody coated beads and receptor were 179 was first chemically conjugated to tosyl-activated prior to adding the polysome library.

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Screening of the various libraries described above yielded the TPO receptor binding peptides shown in Tables 1 and 2 below, as well as others not listed herein.

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× S R N H Δ YSSC L M H M C R R P K LPIHC G A S 'n V L ۳ ı, J U 4 Ŋ X X X e E > œ z U U U Œ, U U Ų U Ö ď Ö U 4 U œ ы × H J LTAWLLE E W I S F H W L D TLSRWLE Ŋ DZ: ß Ŋ > χ Ε r A × > ۲ ω E W L T .1 3 Ö 'J 4 H S 3 3 TABLE A Peptide × Ŋ Z L .] ∡ ü IJ 3 2 H > 3 3 3 3 3 Σ 3 0 REWL GPSLXS E EWL GWL LKEWL GPFWAKAC GLTLREW **≆** 3 o CVMW Ĺ, ы 3 3 ø o . ГЛ s T ш ы œ œ ш 77 24 'J R ∝ ш o ᆆ ם H H H တ o; œ J H 4 œ L u ч G 79 ۲ 7 H H G, ۲ H G T F. H H r R ᆸ .⊐ Eы H H H H > 0 H G G ٢ Δ, Δ, Д, Н H H H G P d, U ρ, o. U α. Ό o. a, Д ۵, Ö O ტ Ö ۵, D, S G о ы o ဗ > ш ... Ö ø o v O U Ω O ш > ω ω O O Ů H C R ø o Δ E a ם ۷ ۲ ۲ D U o ø 4 œ æ Ω Ω n L E E O ш æ ω Ω ы O G Σ > z 4 4 > Ġ S Ø 'n 2 15 20

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WO 96/40750 PCT/US96/09623		EGPTLREWL	T K G P T L R Q W L K S	NAUGPTLROW	T I E G P T L R E W L T	ASDGPTLKEWLSVIRGAS			TABLE 2			S	CRRSELLERC	CTFKQFLDGC	CTRGEWLRCC	CTLRQWLQGC	CTLEELRACC		0	CNRNDLLFC	CIRTEWLHGC	CHLEFMNGC	S	2	H	TRSBWLBRC	CILHEYLSGC	CTREELLRQC	CIFREFUNGC	CSRADFLAAC	SCAQVVOCC	CTLROWILLGMC	
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GSHGCTLREWLCMKIVPC Q W Q G C T L R D C I L R G V F W S SVNSCTLREFLTGCRVFC SYDGCTLRHWLMDIYGDC QRSGCTLRDWVLLNCLAS 29 CTLREFLIMGAC CTLKEWLLWSSC CTLLEWLRNPVC CTLROWLGDAWC CTLGOWLOMGMC CLLLEFLSGADC CTLGEFLAGHLC CRLREFLVDLTC CSFRSWLVDQTC CTLREWLEDIGC CTLODWLVSWTC CTLSEWLSELSC CTLMQWLGGWPC CTLREWLSYGTC CTLQEWLSGGLC CTLREWVFAGLC CTLWGCGKRGC CTRSQWLEGC CSLQEFKHGC CTLGEWKRGC CTLAEFRRGC CTSTOWLLAC CTLREWLEGC CTLOEWRGGC CTRLSGCWLC CTRTOWLLDC CSRSQFLRSC

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IC<sub>50</sub> values for some additional representative peptides are given in the table below. A variety of methods can be used to evaluate IC<sub>50</sub> values. For example, an equilibrium binding ELISA assay, using either MBP-TPO or lacI-peptide tracer, was used to determine whether the peptides inhibit the binding of TPO to the extracellular domain of the TPO receptor. Typically, the IC<sub>50</sub> value were

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C-terminally amidated, or can be prepared as an ester or other determined using the free peptide, which optionally can be The ICSO value can be determined using the free peptide. carboxy amide.

These glycines are not believed to be necessary for binding or activity. Likewise, to mimic the exact sequence of peptides displayed on polysomes, the C-terminal amino acids of the synthetic peptides are often preceded by the sequence M  $\lambda$ Again, this sequence is not believed to be necessary for synthetic peptides are often preceded by one or two glycine To recreate the exact sequence displayed on the phage, the N-terminal and C-terminal amino acids of the binding or activity. residues.

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Those peptides for which ICso values exact sequence displayed by the phage. These glycines are not peptides having the structure: GGCTLREWLHGGFCG were not determined are listed as "N.D.". . The  ${\tt IG_{50}}$  value for gave  ${\sf IC_{S0}}$  values of 500 nm or less are indicated with a "++". Those peptides which gave  $\mathsf{IC}_{\mathsf{SD}}$  values at or near the cutoff G was 500 nm or less. (Note the N-terminal and C-terminal than or equal to 200  $\mu M$  are given a "+", while those which which showed IC $_{50}$  values in excess of 200  $\mu M$  are indicated with a "-". Those peptides which gave  ${\rm IC}_{50}$  values of less point for a particular symbol are indicated with a hybrid symbols "-", "+", and "++". For examples, those peptides amino acids were preceded by two glycines to recreate the  ${\sf IC}_{{\sf SO}}$  values are indicated symbolically by the believed to be necessary for binding or activity.) designator, e.g., "+/-".

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that a preferred core peptide comprises a sequence of amino The tables above, especially Table 3, illustrate

X1 X2 X3 X4 X5 X6 X7

genetically coded L-amino acids;  $X_{\mathsf{S}}$  is A, D, E, G, K, M, Q, R, where X<sub>1</sub> is C, L, M, P, Q, V; X<sub>2</sub> is F, K, L, N, O, R, S, T or S, T, V or Y; X6 is C, F, G, L, M, S, V, W or Y; and X7 is C, V; X3 is C, F, I, L, M, R, S, V or W; X4 is any of the 20 G, I, K, L, M, N, R or V.

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In a preferred embodiment the core peptide comprises a sequence of amino acids:

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X<sub>8</sub> G X<sub>1</sub> X<sub>2</sub> X<sub>3</sub> X<sub>4</sub>X<sub>5</sub> W X<sub>7</sub>

is independently selected from any of the 20 genetically coded Preferred embodiment,  $X_1$  is P;  $X_2$  is T;  $X_3$  is L;  $X_4$  is R;  $X_5$  is where X<sub>1</sub> is L, M, P, Q, or V; X<sub>2</sub> is F, R, S, or T; X<sub>3</sub> is F, L, V, or W; X4 is A, K, L, M, R, S, V, or T; X5 is A, E, G, K, M, Q, R, S, or T; X, is C, I, K, L, M or V; and each Xg residue L-amino acids and their stereoisomeric D-amino acids. In a independently selected from any of the 20 genetically coded non-natural amino acids. Preferably, each  $X_{\rm B}$  residue is L-amino acids, their stereoisomeric D-amino acids; and E or Q; and X, is I or L.

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More preferably, the core peptide comprises a sequence of amino acids:

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More preferably, X9 is where  $X_9$  is A, C, E, G, I, L , M, P, R, Q, S, T, or V; and  $X_9$ X9 X8 G X1 X2 X3 X4 X5 W X7 is A, C, D, E, K, L, Q, R, S, T, or V. A or I; and Xg is D, E, or K.

Particularly preferred peptides include: G G C A D G LKSREHTS; SIE GPTLREWLTSRTPHS; LAIE N, GGCADGPTLREWISFCGGK, TIKGPTLROW PTLREWISFCGG; GNADGPTLRQWLEGRRPK GPTLRQWLHGNGRDT; CADGPTLREWISF and I'E GPTLROWLAARA. 35

peptides for use in this invention include peptides having a In further embodiments of the invention, preferred core structure comprising sequence of amino acids: sequence of amino acids:

C X2 X3 X4 X5 X6 X7

Y; and X; is C, G, I, K, L, M, N, R or V. In a more preferred V. Particularly preferred peptides include: GGCTLREW In a further embodiment,  $X_2$  is S or T;  $X_3$  is L or R;  $X_4$  is R; where X<sub>2</sub> is K, L, N, Q, R, S, T or V; X<sub>3</sub> is C, F, I, L, M, R, S or  $V_i$   $X_4$  is any of the 20 genetically coded L-amino acids; Xs is A, D, E, G, S, V or Y; Xg is C, F, G, L, M, S, V, W or X5 is D, E, or G; X6 is F, L, or W; and X7 is I, K, L, R, or embodiment,  $X_4$  is A, E, G, H, K, L, M, P, Q, R, S, T, or W. LHGGFCGG.

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In a further embodiment, preferred peptides for use in this invention include peptides having a structure comprising a sequence of amino acids:

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XB C X2 X3 X4 X5 X6 X7

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acids; X<sub>5</sub> is A, D, E, G, K, M, Q, R, S, T, V or Y; X<sub>6</sub> is C, F, G, L, M, S, V, W or Y; X, is C, G, I, K, L, M, N, R or V; and  $\chi_{\rm g}$  is any of the 20 genetically coded L-amino acids. In some where X<sub>2</sub> is F, K, L, N, Q, R, S, T or V; X<sub>3</sub> is C, F, I, L, M, R, S, V or W;  $X_4$  is any of the 20 genetically coded L-amino embodiments, X<sub>B</sub> is preferably G, S, Y, or R.

invention. Preferably, for diagnostic purposes, the peptides for pharmaceutical purposes, the peptides and peptidomimetics greater than about 100 mM lack sufficient binding to permit use in either the diagnostic or therapeutic aspects of this and peptidomimetics have an  $IC_{50}$  of about 2 mM or less and, Peptides and peptidomimetics having an  $IC_{50}$  of have an ICso of about 100  $\mu M$  or less.

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system or the "very large scale immobilized polymer synthesis" The binding peptide sequence also provides a means peptide with such activity, but one can also make all of the to determine the minimum size of a TPOR binding compound of the invention. Using the "encoded synthetic library" (ESL) system, one can not only determine the minimum size of a

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peptides that form the group of peptides that differ from the two, or more residues. This collection of peptides can then preferred motif (or the minimum size of that motif) in one, synthesis methods can also be used to synthesize truncation combinations thereof all of the peptide compounds of the immobilized polymers synthesis systems or other peptide be screened for ability to bind to TPO-raceptor. These analogs, deletion analogs, substitution analogs, and invention.

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Example 2 below. Cell proliferation is measured by techniques known in the art, such as an MTT assay which correlates with cell proliferation assay, as described in greater detail in invention were also evaluated in a thrombopoietin dependent TPO-R transfected Ba/F3 cells in a dose dependent manner as The peptides and peptide mimetics of the present (1983)). The peptides tested stimulated proliferation of shown in Figure 1A. These peptides have no effect on the proliferation (see Mossmann J. Immunol. Methods 65:55 H-thymidine incorporation as an indication of cell parental cell line as shown in Figure 1B.

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depicts the effect of carboplatin titration on platelet counts in mice treated with the indicated amounts of carboplatin (in Figures 7 to 9 show the results of a further assay the invention. In this assay mice are made thrombocytopenic peptides of the invention can ameliorate thrombocytopenia in evaluating activity of the peptides and peptide mimetics of amelioration of carboplatin-induced thrombocytopenia on Day The heavy solid lines represent historical data. Figure 8 mg/kg, intraperitoneally (ip) on Day 0). Figure 9 depicts by peptide AF12513 (513). Carboplatin (CBP; 50-125 mg/kg, represent carboplatin-treated groups in three experiments. untreated animals from three experiments. The solid line with carboplatin. Figure 7 depicts typical results when intraperitoneally) was administered on Day 0. AF12513 (1 mg/kg, ip) was given on Days 1-9. These results show the intraperitoneally) on Day 0. The dashed lines represent Balb/C mice are treated with carboplatin (125 mg/kg nouse model.

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invention can be dimerized or oligomerized, thereby increasing TLREWISFCGG was synthesized (GGCADGPTLREW To investigate parental cell lines. Figure 2C shows the results of the assay C-terminally biotinylated analog of the peptide G G C A D G p streptavidin in serum-free HEPES-buffered RPMI at a 4:1 molar streptavidin (SA)) for both the transfected and parental cell lines. Figure 2B shows the results of the assay for the free biotinylated peptide (AF 12285) for both the transfected and I S F C G K (Biotin)). The peptide was preincubated with for streptavidin alone for both the transfected and parental the effect that peptide dimerization/oligomerization has on parental peptide. Figure 2A shows the results of the assay alongside free biotinylated peptide and the unbiotinylated cell lines. These figures illustrate that the pre-formed proliferation of TPO-R transfected Ba/F3 cells, as above, complex was approximately 10 times as potent as the free In addition, certain peptides of the present ratio. The complex was tested for stimulation of cell for the complexed biotinylated peptide (AF 12885 with TPO mimetic potency in cell proliferation assays, a the affinity and/or activity of the compounds. peptide.

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multi-potential primitive haematopoietic progenitor cell line (See, e.g., Dexter et al. <u>J. Exp. Med.</u> 152:1036-1047 (1981)) The specificity of the binding and activity of the peptides of the invention was also examined by studying the This human or murine EPO-R to produce the FDCP-1-EPO-R cell line. This cell line can proliferate, WEHI-3-conditioned media (a medium that contains IL-3, ATCC haematopoietin growth factor receptor family, as is TPO-R. number TIB68). The parental cell line is transfected with The peptides of the invention, as well as TPO, EPO, and a cross reactivity of the peptides for the erythropoieitin assay utilized FDCP-1, a growth factor dependent murine proliferation assay using an EPO-dependent cell line. receptor (EPO-R). The EPO-R is also a member of the known EPO-binding peptide, were examined in a cell but not differentiate when supplemented with as the parental cell line.

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These transfected cell lines can proliferate, but not differentiate in the presence of human or murine EPO.

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The cells were grown to half stationary density in the presence of the necessary growth factors. The cells are then washed in PBS and starved for 16-24 hours in whole media without the growth factors. After determining the viability of the cells, stock solutions (in whole media without the growth factors) are made to give about 10<sup>5</sup> cells per 50 microliters. Serial dilutions of the compounds (typically, the free solution phase peptide as opposed to a phage-bound or other bound or immobilized peptide) to be tested are made in 96-well tissue culture plates for a final volume of 50 microliters per well. Cells (50 microliters) are added to each well and the cells are incubated for 24-48 hours, at which point the negative controls should die or be quiescent. Cell proliferation is then measured by techniques known in the att, such as an WIT assay.

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Figures 3A-G show the results of a series of control proliferation assay using the EPO-dependent cell line. Figure 3F depicts the results for EPO in the cell proliferation assay the results for EPO in the cell proliferation assay using the results for the corresponding parental cell line are shown in Figure 3D. Figure 3E depicts the results for TPO in the cell experiments showing the activity of TPO, the peptides of the present invention, EPO, and EPO-R binding peptides in a cell proliferation assay using either the TPO-R transfected  $\mathtt{Ba/F3}$ EPO-dependent cell line and its corresponding parental line. line and its corresponding parental line. Figure 3B depicts biotinylated peptide (AF 12285 with streptavidin (SA)) and a parental line. Figure 3C depicts the results for complexed proliferation assay using the TPO-R transfected Ba/F3 cell complexed form of a biotinylated EPO-R binding peptide (AF using the EPO-dependent cell line. Figure 3G depicts the results for complexed biotinylated peptide (AF 12285 with 11505 with SA) in the TPO-R transfected Ba/F3 cell line. TPO-R transfected Ba/F3 cell line and its corresponding cell line and its corresponding parental line, or an Figure 3A depicts the results for TPO in the cell

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EPO-R binding peptide (AF 11505 with SA) in the EPO-dependent streptavidin (SA)) and the complexed form of a biotinylated invention bind and activate the TPO-R with a high degree of These results show that the peptides of the specificity. cell line.

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#### PREPARATION OF PETITIES AND PEPTIDE MIMETICS Ä.

#### SOLID PHASE SYNTHESIS

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and even by recombinant DNA technology. See, e.g., Merrifield standard solid phase techniques. The standard methods include exclusive solid phase synthesis, partial solid phase synthesis alpha-amino protected resin. A suitable starting material can Ind. (London) 38:1597 (1966). The benzhydrylamine (BHA) resin methods, fragment condensation, classical solution synthesis, (1970) and is commercially available from Beckman Instruments, resin, or a benzhydrylamine resin. One such chloromethylated alpha-amino acid to a chloromethylated resin, a hydroxymethyl resin is sold under the tradename BIO.BEADS SX-1 by Bio Rad The peptides of the invention can be prepared by has been described by Pietta and Marshall <u>Chem. Commn.</u> 650 classical methods known in the art, for example, by using commenced from the C-terminal end of the peptide using an J. Am. Chem. Soc. 85:2149 (1963), incorporated herein by reference. On solid phase, the synthesis is typically Laboratories, Richmond, CA, and the preparation of the hydroxymethyl resin is described by Bodonszky ec al. be prepared, for instance, by attaching the required Inc., Palo Alto, CA, in the hydrochloride form.

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Thus, the compounds of the invention can be prepared hydrochloric acid (HCl) solutions in organic solvents at room chloromethylated resin with the aid of, for example, cesium Gisin Helv. Chim. Acta. 56:1467 (1973). After the initial bicarbonate catalyst, according to the method described by coupling, the alpha-amino protecting group is removed by a choice of reagents including trifluoroacetic acid (TFA) or by coupling an alpha-amino protected amino acid to the temperature.

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The alpha-amino protecting groups are those known to aliphatic urethane protecting groups (e.g. t-butyloxycarbonyl (Boc), isopropyloxycarbonyl, cyclohexyloxycarbonyl) and alkyl protecting group must be removable upon the completion of the synthesis of the final peptide and under reaction conditions trifluoroacetyl, acetyl), aromatic urethane type protecting type protecting groups (e.g. benzyl, triphenylmethyl). Boc protecting group remains intact during coupling and is not and Fmoc are preferred protecting groups. The side chain groups (e.g. benzyloxycarboyl (Cbz) and substituted Cbz), be useful in the art of stepwise synthesis of peptides. split off during the deprotection of the amino-terminus Included are acyl type protecting groups (e.g. formyl, The side chain that will not alter the target peptide. protecting group or during coupling.

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and 2,5-dichlorobenzyl. The side chain protecting groups for 2,6-dichlorobenzyl, and Cbz. The side chain protecting group for Arg include nitro, Tosyl (Tos), Cbz, adamantyloxycarbonyl tetrahydropyranyl, tert-butyl, trityl, benzyl, Cbz, Z-Br-Cbz, cyclohexyl. The side chain protecting groups for Thr and Ser for Thr and Ser is benzyl. The side chain protecting groups (2-Cl-Cbz), 2-bromobenzyloxycarbonyl (2-BrCbz), Tos, or Boc. include acetyl, benzoyl, trityl, tetrahydropyranyl, benzyl, The side chain protecting groups for Tyr include Asp include benzyl, 2,6-dichlorobenzyl, methyl, ethyl, and mesitoylsulfonyl (Mts), or Boc. The side chain protecting groups for Lys include Cbz, 2-chlorobenzyloxycarbonyl

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the desired order. An excess of each protected amino acid is After removal of the alpha-amino protecting group, the remaining protected amino acids are coupled stepwise in generally used with an appropriate carboxyl group activator example, in methylene chloride  $(CH_2Cl_2)$ , dimethyl formamide such as dicyclohexylcarbodiimide (DCC) in solution, for (DMF) mixtures.

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support by treatment with a reagent such as trifluoroacetic completed, the desired peptide is decoupled from the resin acid or hydrogen fluoride (HF), which not only cleaves the After the desired amino acid sequence has been

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These solid phase peptide synthesis procedures are well known in the art and further described in Stewart <u>Solid</u> Phase Peptide Syntheses (Freeman and Co., San Francisco, (1969)).

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Using the "encoded synthetic library" or "very large scale immobilized polymer synthesis" system described in U.S. Patent Application Serial Nos. 07/492,462, filed March 7, 1990; 07/624,120, filed December 6, 1990; and 07/805,727, filed December 6, 1991; one can not only determine the minimum size of a peptide with such activity, one can also make all of the peptides that form the group of peptides that differ from the preferred motif (or the minimum size of that motif) in one, two, or more residues. This collection of peptides can then be screened for ability to bind to TPO-R. This immobilized polymer synthesis system or other peptide synthesis methods can also be used to synthesize truncation analogs and deletion analogs and combination of truncation and deletion analogs of all of the peptide compounds of the invention.

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#### SYNTHETIC AMINO ACIDS

These procedures can also be used to synthesize peptides in which amino acids other than the 20 naturally occurring, genetically encoded amino acids are substituted at

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one, two, or more positions of any of the compounds of the invention. For instance, naphthylalanine can be substituted for tryptophan, facilitating synthesis. Other synthetic amino acids that can be substituted into the peptides of the present invention include L-hydroxypropyl, L-3, 4-dihydroxyphenylalanyl, d amino acids such as L-d-hydroxyphenylalanyl, b amino acids and non-naturally occurring synthetic amino acids can also be incorporated into the peptides of the present invention.

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One can replace the naturally occurring side chains of the 20 genetically encoded amino acids (or D amino acids) with other side chains, for instance with groups such as alkyl, lower alkyl, cyclic 4-, 5-, 6-, to 7-membered alkyl, amide lower alkyl, amide di(lower alkyl), lower alkoy, hydroxy, carboxy and the lower ester derivatives thereof, and with 4-, 5-, 6-, to 7-membered hetereocyclic. In particular, proline analogs in which the ring size of the proline residue is changed from 5 members to 4, 6, or 7 members can be employed. Cyclic groups can be saturated or unsaturated, and if unsaturated, can be aromatic or non-aromatic.

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Cyclic groups can be saturated or unsaturated, and if unsaturated, can be aromatic or non-aromatic. Heterocyclic groups preferably contain one or more nitrogen, oxygen, and/ox sulphur heteroatoms. Examples of such groups include the fuzazanyl, furyl, imidazolldinyl, imidazollyl, imidazollnyl, oxazolyl, imidazolyl, imidazolinyl, oxazolyl, piperazinyl (e.g. 1-piperazinyl), piperazinyl (e.g. 1-piperidyl, piperazinyl), pyranyl, pyrazinyl, pyrazolidinyl, pyrazolinyl, pyrazolinyl, pyrazolinyl, pyrazolinyl, pyrzolinyl, pyrzolinyl, thiadiazolyl, thienyl, thienyl, thiomorpholinyl (e.g. thiomorpholin), and triazolyl, these heterocyclic groups can be substituted or unsubstituted. Where a group is substituted, the substituted be nusubstituted phenyl.

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One can also readily modify the peptides of the instant invention by phosphorylation, and other methods for

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making peptide derivatives of the compounds of the present invention are described in Hruby et al.<sup>42</sup> Thus, the peptide compounds of the invention also serve as a basis to prepare peptide mimetics with similar biological activity.

The peptide compounds of the invention, including peptidomimetics, can be covalently modified to one or more of a variety of nonproteinaceous polymers, e.g., polyethylene glycol, polypropylene glycol, or polyoxyalkenes, in the manner set forth in U.S. Patent No. 4,640,835; U.S. Patent No. 4,496,689; U.S. Patent No. 4,670,417; U.S. Patent No. 4,791,192; or U.S. Patent No. 4,791,192; or U.S. Patent No. 4,179,337, all which are incorporated by reference in their entirety herein.

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# C. TERMINAL MODIFICATIONS

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of techniques are available for constructing peptide mimetics modifications can be coupled in one peptide mimetic structure inclusion of a -CH2-carbamate linkage between two amino acids Those of skill in the art recognize that a variety group, the C-terminal carboxyl group, and/or changing one or with the same or similar desired biological activity as the stability, and susceptibility to hydrolysis and proteolysis. preparing peptide mimetics modified at the N-terminal amino See, for example, Morgan and Gainor Ann, Rep, Med. Chem. (e.g., modification at the C-terminal carboxyl group and more of the amido linkages in the peptide to a non-amido 24:243-252 (1989). The following describes methods for corresponding peptide compound but with more favorable activity than the peptide with respect to solubility, linkage. It being understood that two or more such in the peptide).

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## 1. N-TERMINAL MODIFICATIONS

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The peptides typically are synthesized as the free acid but, as noted above, could be readily prepared as the amide or ester. One can also modify the amino and/or carboxy

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terminus of the peptide compounds of the invention to produce other compounds of the invention. Amino terminus modifications include methylating (i.e., 'NHCH<sub>3</sub> or 'NH(CH<sub>3</sub>)<sub>2</sub>), acetylating, adding a carbobenzoyl group, or blocking the amino terminus with any blocking group containing a carboxylate functionality defined by RCOO-, where R is selected from the group consisting of naphthyl, acridinyl, steroidyl, and similar groups. Carboxy terminus modifications include replacing the free acid with a carboxamide group or forming a cyclic lactam

at the carboxy terminus to introduce structural constraints.

Amino terminus modifications are as recited above and include alkylating, acetylating, adding a carbobenzoyl group, forming a succinimide group, etc. Specifically, the N-terminal amino group can then be reacted as follows:

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(a) to form an amide group of the formula RC(O)NH-where R is as defined above by reaction with an acid halide [e.g., RC(O)Cl] or acid anhydride. Typically, the reaction can be conducted by contacting about equimolar or excess amounts (e.g., about 5 equivalents) of an acid halide to the peptide in an inert diluent (e.g., dichloromethane) preferably containing an excess (e.g., about 10 equivalents) of a tertiary amine, such as diisopropylethylamine, to scavenge the acid generated during reaction. Reaction conditions are otherwise conventional (e.g., room temperature for 30 minutes). Alkylation of the terminal amino to provide for a lower alkyl N-substitution followed by reaction with an acid halide as described above will provide for N-alkyl amide group of the formula RC(O)NR-;

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(b) to form a succinimide group by reaction with succinic anhydride. As before, an approximately equimolar amount or an excess of succinic anhydride (e.g., about 5 equivalents) can be employed and the amino group is converted to the succinimide by methods well known in the art including the use of an excess (e.g., ten equivalents) of a tertiary amine such as disopropylethylamine in a suitable inert solvent (e.g., dichloromethane). See, for example, Wollenberg, et al., U.S. Patent No. 4,612,132 which is incorporated herein by reference in its entirety. It is

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understood that the succinic group can be substituted with, for example, C<sub>2</sub>-C<sub>6</sub> alkyl or -SR substituents which are prepared in a conventional manner to provide for substituted succinimide at the N-terminus of the peptide. Such alkyl substituents are prepared by reaction of a lower olefin (C<sub>2</sub>-C<sub>6</sub>) with maleic anhydride in the manner described by Wollenberg, et al., supra. and -SR substituents are prepared by reaction of RSH with maleic anhydride where R is as defined above;

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lenzyloxycarbonyl-NH- or a substituted benzyloxycarbonyl-NH- group by reaction with approximately an equivalent amount or an excess of CBZ-Cl (i.e., benzyloxycarbonyl chloride) or a substituted CBZ-Cl in a sultable inert diluent (e.g., dichloromethane) preferably containing a tertiary amine to scavenge the acid generated during the reaction,

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(d) to form a sulfonamide group by reaction with an equivalent amount or an excess (e.g., 5 equivalents) of R-S(0)<sub>2</sub>Cl in a suitable inert diluent (dichloromethane) to convert the terminal amine into a sulfonamide where R is as defined above. Preferably, the inert diluent contains excess tertiary amine (e.g., ten equivalents) such as disopropylethylamine, to scavenge the acid generated during reaction. Reaction conditions are otherwise conventional (e.g., room temperature for 30 minutes);

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(e) to form a carbamate group by reaction with an equivalent amount or an excess (e.g., 5 equivalents) of R-OC(O)Cl or R-OC(O)Oc<sub>6</sub>H<sub>4</sub>-p-NO<sub>2</sub> in a suitable inert diluent (e.g., dichloromethane) to convert the terminal amine into a carbamate where R is as defined above. Preferably, the inert diluent contains an excess (e.g., about 10 equivalents) of a tertiary amine, such as diisopropylethylamine, to scavenge any acid generated during reaction. Reaction conditions are otherwise conventional (e.g., room temperature for 30 minutes); and

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(f) to form a urea group by reaction with an equivalent amount or an excess (e.g., 5 equivalents) of R-N=C=O in a suitable inert diluent (e.g., dichloromethane) to

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convert the terminal amine into a urea (1.e., RNHC(0)NH-) group where R is as defined above. Preferably, the inert diluent contains an excess (e.g., about 10 equivalents) of a tertiary amine, such as dissopropylethylamine. Reaction conditions are otherwise conventional (e.g., room temperature for about 30 minutes).

## . C-TERMINAL MODIFICATIONS

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In preparing peptide mimetics wherein the C-terminal carboxyl group is replaced by an ester (i.e., -C(0)OR where R is as defined above), the resins used to prepare the peptide acids are employed, and the side chain protected peptide is cleaved with base and the appropriate alcohol, e.g., methanol. Side chain protecting groups are then removed in the usual fashion by treatment with hydrogen fluoride to obtain the desired ester.

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benzhydrylamine resin is used as the solid support for peptide In preparing peptide mimetics wherein the C-terminal Side chain protection is then removed in the usual fashion by chloromethylated resin during peptide synthesis coupled with C-terminus is -C(0)NRR $^{1}$  where R and R $^{1}$  are as defined above). reaction with an alkylamine or a dialkylamine yields a side fluoride treatment to release the peptide from the support peptide from the support yields the free peptide amide and treatment with hydrogen fluoride to give the free amides, reaction with ammonia to cleave the side chain protected synthesis. Upon completion of the synthesis, hydrogen carboxyl group is replaced by the amide -C(0) NR<sup>3</sup>R<sup>4</sup>, a results directly in the free peptide amide (i.e., the chain protected alkylamide or dialkylamide (i.e., the C-terminus is -C(0)NH2). Alternatively, use of the alkylamídes, or dialkylamides.

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In another alternative embodiment, the C-terminal carboxyl group or a C-terminal ester can be induced to cyclize by internal displacement of the -OH or the ester (-OR) of the carboxyl group or ester respectively with the N-terminal amino group to form a cyclic peptide. For example, after synthasis

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dimethyl formamide (DMF) mixtures: The cyclic peptide is then formed by internal displacement of the activated ester with converted to an activated ester by an appropriate carboxyl group activator such as dicyclohexylcarbodiimide (DCC) in the N-terminal amine. Internal cyclization as opposed to and cleavage to give the peptide acid, the free acid is solution, for example, in methylene chloride  $(CH_2Cl_2)$ , polymerization can be enhanced by use of very dilute Such methods are well known in the art. solutions.

terminii of the peptide, so that there is no terminal amino or include amide, amide lower alkyl, amide di(lower alkyl), lower carboxyl group, to decrease susceptibility to proteases or to alkoxy, hydroxy, and carboxy, and the lower ester derivatives One can also cyclize the peptides of the invention, functional groups of the compounds of the present invention thereof, and the pharmaceutically acceptable salts thereof. or incorporate a desamino or descarboxy residue at the C-terminal restrict the conformation of the peptide.

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#### BACKBONE MODIFICATIONS ä

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biological activity as the lead peptide compound but with more See Morgan and Gainor <u>Ann. Rep. Med. Chem.</u> 24:243-252 (1989), al. Biochem J. 268(2):249-262 (1990), incorporated herein by reference. Thus, the peptide compounds of the invention also Other methods for making peptide derivatives of the compounds of the present invention are described in Hruby et favorable activity than the lead with respect to solubility, stability, and susceptibility to hydrolysis and proteolysis. incorporated herein by reference. These techniques include phosphonates, amidates, carbamates, sulfonamides, secondary serve as structural models for non-peptidic compounds with replacing the peptide backbone with a backbone composed of recognize that a variety of techniques are available for similar biological activity. Those of skill in the art constructing compounds with the same or similar desired umines, and N-methylamino acids.

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Peptide mimetics wherein one or more of the peptidyl linkages [-C(O)NH-] have been replaced by such linkages as a  $-CH_2$ -sulfonamide linkage, a urea linkage, a secondary amine suitably protected amino acid analogue for the amino acid [-C(O)NR6- where R6 is lower alkyl] are prepared during conventional peptide synthesis by merely substituting a (-CH2NH-) linkage, and an alkylated peptidyl linkage reagent at the appropriate point during synthesis. -CH $_2$ -carbamate linkage, a phosphonate linkage, a

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with the free amine or an alkylated amine on the N-terminus of (-CH<sub>2</sub>OC(O)NR-), then the carboxyl (-COOH) group of a suitably leads to the formation of a -CH2OC(O)NR- linkage. For a more -C(0)NR- linkage in the peptide with a -CH2-carbamate linkage which is then converted by conventional methods to a -OC(0)Cl functionality. Reaction of either of such functional groups Suitable reagents include, for example, amino acid detailed description of the formation of such -CH $_2$ -carbamate been replaced with a moiety suitable for forming one of the the partially fabricated peptide found on the solid support analogues wherein the carboxyl group of the amino acid has above linkages. For example, if one desires to replace a protected amino acid is first reduced to the -CH2OH group functionality or a para-nitrocarbonate -OC(0)0-C<sub>6</sub>H<sub>4</sub>-p-NO<sub>2</sub> linkages, see Cho et al <u>Science</u>, 261:1303-1305 (1993).

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which are incorporated herein by reference in their entirety. Similarly, replacement of an amido linkage in the 07/943,805, 08/081,577, and 08/119,700, the disclosures of peptide with a phosphonate linkage can be achieved in the manner set forth in U.S. Patent Application Serial Nos.

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Replacement of an amido linkage in the peptide with example, thioacetic acid followed by hydrolysis and oxidative carboxyl (-COOH) group of a suitably protected amino acid to the -CH2OH group and the hydroxyl group is then converted to suitable leaving group such as a tosyl group by conventional a -CH2-sulfonamide linkage can be achieved by reducing the chlorination will provide for the -CH2-S(0)2Cl functional methods. Reaction of the tosylated derivative with, for group which replaces the carboxyl group of the otherwise

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Suitably protected amino acid. Use of this suitably protected amino acid analogue in peptide synthesis provides for inclusion of an -CH<sub>2</sub>S(0)<sub>2</sub>NR- linkage which replaces the amido linkage in the peptide thereby providing a peptide mimetic. For a more complete description on the conversion of the carboxyl group of the amino acid to a -CH<sub>2</sub>S(0)<sub>2</sub>Cl group, see, for example, Weinstein, Boris Chemistry & Biochemistry of Amino Acids, Peptides and Proteins Vol. 7, pp. 267-157, Marcel Dekker, Inc., New York (1981) which is incorporated herein by reference.

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Replacement of an amido linkage in the peptide with a urea linkage can be achieved in the manner set forth in U.S. Patent Application Serial No. 08/147,805 which application is incorporated herein by reference in its entirety.

Secondary amine linkages wherein a -CH<sub>2</sub>NH- linkage replaces the amido linkage in the peptide can be prepared by employing, for example, a suitably protected dipeptide analogue wherein the carbonyl bond of the amido linkage has been reduced to a CH<sub>2</sub> group by conventional methods. For example, in the case of diglycine, reduction of the amide to the amine will yield after deprotection H<sub>2</sub>NCH<sub>2</sub>CH<sub>2</sub>NHCH<sub>2</sub>COOH which is then used in N-protected form in the next coupling reaction. The preparation of such analogues by reduction of the carbonyl group of the amido linkage in the dipeptide is well known in the art.

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The suitably protected amino acid analogue is employed in the conventional peptide synthesis in the same manner as would the corresponding amino acid. For example, typically about 3 equivalents of the protected amino acid analogue are employed in this reaction. An inert organic diluent such as methylene chloride or DMF is employed and, when an acid is generated as a reaction by-product, the reaction solvent will typically contain an excess amount of a tertiary amine to scavenge the acid generated during the reaction. One particularly preferred tertiary amine is disopropylethylamine which is typically employed in about 10 fold excess. The reaction results in incorporation into the peptide mimetic of an amino acid analogue having a

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non-peptidyl linkage. Such substitution can be repeated as desired such that from zero to all of the amido bonds in the peptide have been replaced by non-amido bonds.

One can also cyclize the peptides of the invention, or incorporate a desamino or descarboxy residue at the termini of the peptide, so that there is no terminal amino or carboxyl group, to decrease susceptibility to proceases or to restrict the conformation of the peptide. C-terminal functional groups of the compounds of the present invention include amide, amide lower alkyl, amide di(lower alkyl), lower alkoxy, hydroxy, and carboxy, and the lower ester derivatives thereof, and the pharmaceutically acceptable salts thereof. Examples of cyclized compounds are provided in Tables 4, 5, 6, 8, and 9.

## E. DISULFIDE BOND FORMATION

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The compounds of the present invention may exist in a cyclized form with an intramolecular disulfide bond between the thiol groups of the cysteines. Alternatively, an intermolecular disulfide bond between the thiol groups of the cysteines can be produced to yield a dimeric (or higher oligomeric) compound. One or more of the cysteine residues may also be substituted with a homocysteine. These intramolecular or intermolecular disulfide derivatives can be represented schematically as shown below:

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wherein m and n are independently 1 or 2.

Other embodiments of this invention provide for analogs of these disulfide derivatives in which one of the

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sulfurs has been replaced by a CH<sub>2</sub> group or other isostere for sulfur. These analogs can be made via an intramolecular or intermolecular displacement, using methods known in the art as shown below:

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wherein p is 1 or 2. One of skill in the art will readily appreciate that this displacement can also occur using other homologs of the a-emino-g-butyric acid derivative shown above and homocysteine.

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Alternatively, the amino-terminus of the peptide can be capped with an alpha-substituted acetic acid, wherein the alpha substituent is a leaving group, such as an a-haloacetic acid, for example, a-chloroacetic acid, a-bromoacetic acid, or a-iodoacetic acid. The compounds of the present invention can be cyclized or dimerized via displacement of the leaving group by the sulfur of the cysteine or homocysteine residue. See, e.g., Barker et al. J. Med. Chem. 35:2040-2048 (1992) and Or et al. J. Org. Chem. 56:3146-3149 (1991), each of which is incorporated herein by reference. Examples of dimerized compounds are provided in Tables 7, 9, and 10.

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#### V. UTILITY

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The compounds of the invention are useful in vitro as unique tools for understanding the biological role of TPO, including the evaluation of the many factors thought to influence, and be influenced by, the production of TPO and the

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receptor binding process. The present compounds are also useful in the development of other compounds that bind to and activate the TPO-R, because the present compounds provide important information on the relationship between structure and activity that should facilitate such development.

The compounds are also useful as competitive binders for example, by labeling, such as covalently or non-covalently assay embodiments, the compounds of the invention can be used without modification or can be modified in a variety of ways; labeling include biotinylation of one constituent followed by labels (U.S. Patent Np. 3,940,475) capable of monitoring the in assays to screen for new TPO receptor agonists. In such such as peroxidase and alkaline phosphatase, and fluorescent Possibilities for direct labeling include label groups such as: radiolabels such as <sup>125</sup>I, enzymes (US Patent 3,645,090) binding to avidin coupled to one of the above label groups. The compounds may also include spacers or linkers in cases where the compounds are to be attached to a solid support. detectable signal. In any of these assays, the materials joining a moiety which directly or indirectly provides a change in fluorescence intensity, wavelength shift, or fluorescence polarization. Possibilities for indirect thereto can be labeled either directly or indirectly.

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Moreover, based on their ability to bind to the TPO receptor, the peptides of the present invention can be used as reagents for detecting TPO receptors on living cells, fixed cells, in biological fluids, in tissue homogenates, in purified, natural biological materials, etc. Por example, by labelling such peptides, one can identify cells having TPO-R on their surfaces. In addition, based on their ability to bind the TPO receptor, the peptides of the present invention can be used in in situ staining, FACS (fluorescence-activated cell sorting), Western blotting, ELISA, etc. In addition, based on their ability to bind to the TPO receptor, the peptides of the present invention can be used in receptor purification, or in purifying cells expressing TPO receptors on the cell surface (or inside permeabilized cells).

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diagnostic applications wherein the TPO-receptor is preferably The compounds of the present invention can also be utilized as commercial reagents for various medical research functional assays; (2) use to maintain the proliferation and transduction/receptor activation; and (5) other research and analysis of the TPO-receptor through co-crystallization; (4) and diagnostic uses. Such uses include but are not limited (1) use as a calibration standard for quantitating the growth of TPO-dependent cell lines; (3) use in structural against a known quantity of a TPO agonist, and the like. activated or such activation is conveniently calibrated activities of candidate TPO agonists in a variety of use to investigate the mechanism of TPO signal

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thrombocycopenia by killing the rapidly dividing, more mature infusing patients post chemotherapy or radiation therapy with The compounds of the present invention can be used Thus, amelioration of the thrombocytopenia by TPO or megakaryocytes and immature precursors by in vitro culture. committed progenitors, both in conjunction with additional cytokines or on their own. See, e.g., DiGiusto et al. PCT treatments can also reduce the number and viability of the Publication No. 95/05843, which is incorporated herein by population of megakaryocytes. However, these therapeutic immature, less mitotically active megakaryocyte precursor the compounds of the present invention can be hastened by for the in vitro expansion of megakaryocytes and their reference. Chemotherapy and radiation therapies cause a population of his or her own cells enriched for cells.

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TPO-R in vivo. For example, the peptides and compounds of the hematological disorders, including but not limited to platelet invention in amounts sufficient to mimic the effect of TPO on encompasses methods for therapeutic treatment of TPO related disorders and thrombocytopenia, particularly when associated administered to warm blooded animals, including humans, to activate the TPO-R in vivo. Thus, the present invention disorders that comprise administering a compound of the The compounds of the invention can also be invention can be administered to treat a variety of

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with bone marrow transfusions, radiation therapy, and

chemotherapy

antagonists are preferably first administered to patients undergoing chemotherapy or radiation therapy, followed by administration of the tpo agonists of the invention. In some embodiments of the invention, TPO

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invention can be evaluated either in vitro or in vivo in one of the numerous models described in McDonald Am. J. of The activity of the compounds of the present Pedlatric Hematology/Oncology 14:8-21 (1992), which is incorporated herein by reference.

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According to one embodiment, the compositions of the or chemotherapy. The compounds typically will be administered prophylactically prior to chemotherapy, radiation therapy, or associated with bone marrow transfusions, radiation therapy, present invention are useful for treating thrombocytopenia bone marrow transplant or after such exposure.

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nasal, vaginal, rectal, or sublingual routes of administration of the invention in association with a pharmaceutical carrier ingredient, at least one of the peptides or peptide mimetics intraperitoneal, intravenous (IV) or subcutaneous injection), Publication No. WO 94/17784; and Pitt et al. European Patent Application 613,683, each of which is incorporated herein by Accordingly, the present invention also provides and can be formulated in dosage forms appropriate for each Patent Publication No. WO 93/25221; Pitt et al. PCT Patent See, e.g., Bernstein et al. PCT administered by oral, pulmonary, parental (intramuscular, inhalation (via a fine powder formulation), transdermal, pharmaceutical compositions comprising, as an active or diluent. The compounds of this invention can be route of administration. reference.

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Solid dosage forms for oral administration include comprise, as is normal practice, additional substances other least one inert pharmaceutically acceptable carrier such as solid dosage forms, the active compound is admixed with at In such sucrose, lactose, or starch. Such dosage forms can also capsules, tablets, pills, powders, and granules.

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pharmaceutically acceptable emulsions, solutions, suspensions, Liquid dosage forms for oral administration include used in the art, such as water. Besides such inert diluents, syrups, with the elixirs containing inert diluents commonly agents, emulsifying and suspending agents, and sweetening, compositions can also include adjuvants, such as wetting flavoring, and perfuming agents.

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They parental administration include sterile aqueous or non-aqueous solutions, suspensions, or emulsions. Examples of non-aqueous bacteria retaining filter, by incorporating sterilizing agents heating the compositions. They can also be manufactured using into the compositions, by irradiating the compositions, or by gelatin, and injectable organic esters such as ethyl oleate. preserving, wetting, emulsifying, and dispersing agents. may be sterilized by, for example, filtration through a solvents or vehicles are propylene glycol, polyethylene sterile water, or some other sterile injectable medium, glycol, vegetable oils, such as olive oil and corn oil, Preparations according to this invention for Such dosage forms may also contain adjuvants such as immediately before use.

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are preferably suppositories which may contain, in addition to administration are also prepared with standard excipients well Compositions for rectal or vaginal administration the active substance, excipients such as cocoa butter or a suppository wax. Compositions for nasal or sublingual known in the art.

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In therapeutic applications, compositions are administered to irrest the symptoms of the disease and its complications. An above, in an amount sufficient to cure or at least partially administered for prophylactic and/or therapeutic treatments. The compositions containing the compounds can be a patient already suffering from a disease, as described

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"therapeutically effective dose". Amounts effective for this use will depend on the severity of the disease and the weight amount adequate to accomplish this is defined as and general state of the patient.

microencapsulated by, for example, the method of Tice and Bibi (in Treatise on Controlled Drug Delivery, ed. A. Kydonieus, The compositions of the invention can also be Marcel Dekker, N.Y. (1992), pp. 315-339).

containing the compounds of the invention are administered to a patient susceptible to or otherwise at risk of a particular disease. Such an amount is defined to be a "prophylactically effective dose". In this use, the precise amounts again In prophylactic applications, compositions depend on the patient's state of health and weight.

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state of the patient, and other medicants administered. Thus, efficacy. Typically, dosages used in vitro may provide useful Various considerations Gilman's: The Pharmacological Basis of Therapeutics, 8th ed., including means of administration, target site, physiological Sciences, 7th ed., Mack Publishing Co., Easton, Penn. (1985); guidance in the amounts useful for in situ administration of treatment dosages should be titrated to optimize safety and effective therapy will depend upon many different factors, The quantities of the TPO agonist necessary for are described, e.g., in Gilman et al. (eds), Goodman and these reagents. Animal testing of effective doses for treatment of particular disorders will provide further Pergamon Press (1990); and Remington's Pharmaceutical each of which is hereby incorporated by reference. predictive indication of human dosage.

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administered at a dosage range of from about 0.001 mg to about The peptides and peptide mimetics of this invention 10 mg/kg of body weight per day. The specific dose employed is regulated by the particular condition being treated, the severity of the condition, the age and general condition of route of administration as well as by the judgement of the are effective in treating TPO mediated conditions when attending clinician depending upon factors such as the the parient, and the like.

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Although only preferred embodiments of the invention are specifically described above, it will be appreciated that modifications and variations of the invention are possible without departing from the spirit and intended scope of the invention.

#### EXAMPLE 1

# SOLID PHASE PEPTIDE SYNTHESIS

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Various peptides of the invention were synthesized using the Merrifield solid phase synthesis techniques (See Steward and Young, <u>Solid Phase Peptide Synthesis</u>, 2d. edition, Pierce Chemical, Rockford, IL (1984) and Merrifield <u>J. Am. Chem. Soc.</u> 85:2149 (1963)) on a Milligen/Biosearch 9600 automated instrument or an Applied Biosystems Inc. Model 431A peptide synthesizer. The peptides were assembled using standard protocols of the Applied Biosystems Inc. System Software version 1.01. Each coupling was performed for mone-two hours with BOP (benzotriazoly)

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N-oxtrisdimethylaminophosphonium hexafluorophosphate) and HOBt (1-hydroxybenzotriazole).

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The resin used was HMP resin or PAL (Milligen/Biosearch), which is a cross-linked polystyrene resin with 5-(4'-Fmoc-aminomethyl-3,5'-dimethyoxyphenoxy) valeric acid as a linker. Use of PAL resin results in a carboxyl terminal amide functionality upon cleavage of the peptide from the resin. Upon cleavage, the HMP resin produces a carboxylic acid moiety at the C-terminus of the final product. Most reagents, resins, and protected amino acids (free or on the resin) were purchased from Millipore or Applied Biosystems Inc.

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The Fmoc group was used for amino protection during the coupling procedure. Primary amine protection on amino acids was achieved with Fmoc and side chain protection groups were t-butyl for serine, tyrosine, asparagine, glutamic acid, and threonine; trityl for glutamine; Pmc (2,2,5,7,8-pentamethylchroma sulfonate) for arginine;

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N-t-butyloxycarbonyl for tryptophan; N-trityl for histidine and glutamine; and S-trityl for cysteine.

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In all cases, achieved by treatment with reagent K or slight modifications performance liquid chromatography on a  $C_{16}$  bonded silica gel acid, 5% ethanedithiol, and 5% water, initially at 4°C, and characterized by Fast Atom Bombardment mass spectrometry or electrospray mass spectrometry and amino acid analysis when simultaneous deprotection of the side chain functions were gradually increasing to room temperature. The deprotected of it. Alternatively, in the synthesis of those peptides, peptide was cleaved with a mixture of 90% trifluoroacetic with an amidated carboxyl terminus, the fully assembled Removal of the peptides from the resin and column with a gradient of acetonitrile/water in 0.1% trifluoroacetic acid. The homogeneous peptides were purification was by preparative, reverse-phase, peptides were precipitated with diethyl ether. applicable.

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#### EXAMPLE 2 BIOASSAYS

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Bioactivity of the peptides can be measured using a thrombopoietin dependent cell proliferation assay. Murine IL-3 dependent Ba/F3 cells were transfected with full length human TPO-R. In the absence of IL-3 (WEHI-3 conditioned media), these cells are dependent on TPO for proliferation. The parental, untransfected cell line does not respond to human TPO, but remains IL-3 dependent.

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Bioassays have been performed on both of the above cell lines using synthetic peptides derived from library screening. The cells were grown in complete RPMI-10 media, containing 10% WEHI-3 conditioned media, then washed twice in PBS, resuspended in media which lacked WEHI-3 conditioned media, and added to wells containing dilutions of peptide or TPO at 2 x 10% cells/well. The cells were incubated for 48 hours at 37°C in a humidified 5% CO<sub>2</sub> atmosphere and metabolic activity was assayed by the reduction of MTT to formazan, with

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absorbance at 570 nM measured on an ELLSA plate reader. The peptides tested stimulated proliferation of TPO-R transfected Ba/F3 cells in a dose dependent manner as shown in Figure 1. These peptides have no effect on the parental cell line.

#### EXAMPLE 3 BINDING AFFINITY

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harvest. Various concentrations of peptide or peptide mimetic mixtures were added to the TPO-R coated wells, incubated for 2 wells were then treated with a 1:10 dilution of soluble TPO-R assay. The wells of a microtiter plate were coated with 1 mg were mixed with a constant amount of a truncated form of TPO maltose binding protein (MBP-TPO $_{156}$ ). The peptide MBP-TPO $_{156}$ streptavidin, blocked with PBS/1% BSA, followed by 50 ng of adding a rabbit anti-sera directed against MBP, followed by alkaline phosphatase conjugated goat anti-rabbit IgG. The peptides for TPO-R were measured in a competition binding biotinylated anti-receptor immobilizing antibody (Ab179). MBP-TPO<sub>156</sub> that was bound at equilibrium was measured by consisting of residues 1-156 fused to the C-terminus of Binding affinities of chemically synthesized hours at 4°C and then washed with PBS. The amount of amount of alkaline phosphatase in each well was then determined using standard methods.

The assay is conducted over a range of peptide concentrations and the results are graphed such that the y axis represents the amount of bound MBP-TPO<sub>156</sub> and the x axis represents the concentration of peptide or peptide mimetic. One can then determine the concentration at which the peptide or peptide mimetic will reduce by 50% (IC<sub>50</sub>) the amount of MBP-TPO<sub>156</sub> bound to immobilized TPO-R. The dissociation constant (Kd) for the peptide should be similar to the measured IC<sub>50</sub> using the assay conditions described above.

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#### EXAMPLE 4 "PEPTIDES ON PLASMIDS"

The pJS142 vector is used for library construction and is shown in Figure 4. Three oligonucleotide sequences are needed for library construction: ON-829 (5' ACC ACC TCC GG); ON-830 (5' TTA CTT AGT TA) and a library specific oligonucleotide of interest (5' GA GGT GGT  $\{NNK\}_n$  TAA CTA AGT AAA GC), where  $\{NNK\}_n$  denotes a random region of the desired length and sequence. The oligonucleotides can be 5' phosphorylated chemically during synthesis or after purification with polynucleotide kinase. They are then annealed at a 1:1:1 molar ratio and ligated to the vector.

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The strain of E. coli which is preferably used for panning has the genotype:  $\Delta(ax1-xecA)$  endAl nupG lon-11 sulAl hsdR17  $\Delta(cmpT-fepC)$ 266  $\Delta clpA319::kan$   $\Delta lacI$  lac ZU18 which can be prepared from an E. coli strain from the E. coli Genetic Stock Center at Yale University (E. coli b/r, stock center designation CGSC:6573) with genotype lon-11 sulAl. The above E. coli strain is prepared for use in electroporation as described by Dower et al. Nucleic Acids Res. 16:6127 (1988), except that 10% glycerol is used for all wash steps. The cells are tested for efficiency using 1 pg of a Bluescript plasmid (Stratagene). These cells are used for growth of the original library and for amplification of the enriched population after each round of panning.

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Peptides on plasmids are released from cells for panning by gentle enzymatic digestion of the cell wall using lysozyme. After pelleting of the cell debris, the crude lysate can be used directly on most receptors. If some additional purification of the plasmid complexes is needed, a gel filtration column can be used to remove many of the low molecular weight contaminants in the crude lysate.

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Panning is carried out in a buffer (HEKL) of a lower salt concentration than most physiological buffers. The panning can be conducted in microtiter wells with a receptor immobilized on a nonblocking monoclonal antibody (MAb) or by panning on beads or on columns. More specifically, in the

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receptor (PAN sample) and 6 wells without receptor (NC sample) are typically used. Comparison of the number of plasmids in these two samples can give an indication of whether receptor first round of panning, 24 wells, each coated with receptor, specific clones are being enriched by panning. "Enrichment" can be used. For the second round, six wells coated with recovered from the NC sample. Enrichment of 10 fold is usually an indication that receptor specific clones are is defined as the ratio of PAN transformants to those present.

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usually 100  $\mu$ l of lysate per well is used. In round 3, 100  $\mu$ l of lysace per well diluted with 1/10 in HEXL/BSA is used. For In later rounds of panning, it is useful to reduce transforming units of at least 1000 fold above the estimated background binding of the plasmid complexes. In round 2, further rounds of panning, typically an input of plasmid the input of lysate into the wells to lower nonspecific remaining diversity is used.

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observed. Typically, an ELISA that detects receptor specific but is a disadvantage in that the signal in the ELISA is not The binding properties of the peptides encoded by cooperative, multivalent fashion. This cooperative binding The sensitivity of this assay is an advantage in sufficient density of receptor can be immobilized in wells, normally a tetramer and the minimum functional DNA binding that initial hits of low affinity can be easily identified, individual clones are typically examined after 3, 4, or 5 Fusion of the peptides to maltose binding protein (MBP) as binding by LacI-peptide fusion proteins is used. LacI is permits the detection of binding events of low intrinsic the peptides fused to LacI will bind to the surface in a described below permits testing in an ELISA format where rounds of panning, depending on the enrichment numbers correlated with the intrinsic affinity of the peptides. multivalently on the fusion protein. Assuming that a The peptides are thus displayed signal strength is better correlated with affinity. species is a dimer. See Figure 5A-B. affinity.

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those sequences in frame with the gene encoding MBP, a protein of a library into pJS142 creates a BspEI restriction site near for ligation. In addition, correct ligation of the Scal sites populations of clones can be transferred to vectors that fuse respectively, available commercially from New England Biolabs. The cloning See Figure 5A-B. Digestion of pELM3 and pELM15 with Agel and Scal allows efficient cloning of the BspEI-Scal fragment from promoter-driven MBP-peptide fusions can then be induced with the pJS142 library. The BspEI and Agel ends are compatible double stranded form using any standard miniprep procedure. vectors, pELM3 (cytoplasmic) or pELM15 (periplasmic), which Digestion with BspEI and Scal allows the purification of a 7900 bp DNA fragment that can be subcloned into one of two are simple modifications of the pMALc2 and pMALp2 vectors, is essential to recreate a functional bla (Amp resistance) the beginning of the random coding region of the library. DNA from clones of interest can be prepared in gene, thus lowering the level of background clones from The coding sequences of interesting single clones or undesired ligation events. Expression of the tac that generally occurs as a monomer in solution.

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Lysates for the LacI or MBP ELISAs are prepared from insoluble cell debris by centrifugation. The lysates are then individual clones by lysing cells using lysozyme and removing added to wells containing immobilized receptor and to control wells without receptor. Binding by the LacI or MBP peptide fusions is detected by incubation with a rabbit polyclonal

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incubation with alkaline phosphatase labeled goat anti rabbit second antibody. The bound alkaline phosphatase is detected antiserum directed against either LacI or MBP followed by with p-nitrophenyl phosphate chromagenic substrate.

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#### EXAMPLE 5

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#### "HEADPIECE DIMER" SYSTEM

A variant of the LacI peptides-on-plasmids technique utilizes a DNA binding protein called "headpiece dimer". DNA approximately 60 amino acid "headpiece" domain. The dimer of of the headpiece dimer with the plasmid encoding that peptide. containing two headpieces connected via short peptide linker. association of a peptide epitope displayed at the C-terminus "headpiece dimer" system utilizes headpiece dimer molecules These proteins bind DNA with sufficient stability to allow binding by the E. coli lac repressor is mediated by the the headpiece domains that binds to the lac operator is approximately 300 amino acid C-terminal domain. The normally formed by association of the much larger

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it to make a peptide-headpiece dimer-plasmid complex that can The random peptides are fused to the C-terminus of the headpiece dimer, which binds to the plasmid that encoded libraries based on initial low-affinity hits, and selecting peptides-on-plasmids system allows greater selectivity for headpiece dimer system is useful for making mutagenesis high affinity ligands than the LacI system. Thus, the higher affinity variants of those initial sequences. be screened by panning. The headpiece dimer

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The libraries performed with aqueous phenol instead of with IPTG. Sequences for LacI libraries, except that HEK buffer is used instead of libraries is carried out by similar procedures to those used 6A-C) . The presence of the lac operator is not required for ZU118 A(srl-recA) 306::Tnl0 .and amplified under conditions of basal (A) promoter induction. Panning of headpiece dimer transfer to the MBP vector so that they can be tested in the The libraries are constructed as with peptides on (lon-11 sulAl hsdRl7 (ompT-fepC) AclpA319::kan AlacI lac from headpiece dimer panning are often characterized after were introduced into bacterial strain comprising  $E.\ coli$ plasmids using headpiece dimer vector pCMG14 (see Figure HEKL buffer and elution of plasmids from the wells is plasmid binding by the headpiece dimer protein.

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clones can be screened by colony lifts with labeled receptor. affinity sensitive MBP ELISA and also so that populations of

#### EXAMPLE 6

In this example cyclized compounds were subjected to determined. The ranges for  $EC_{S_0}$  are symbolically indicated as for  $\mathsf{IC}_{50}$  described above. The results are summarized in Table acidification of the extracellular medium in response to TPO three assays. First, IC50 valves were obtained as described Finally, a microphysiometer (Molecular Devices Corp.) assay receptor stimulation by the compounds of the invention was above. Additionally, an MTT cell proliferation assay as described above was performed to calculate  $EC_{50}$  values. was performed. Basically, in this assay the rate of

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ECSO(nM) ECSO(nM) (CSO(nM) Proliferanon Mercepays. [H] - (D-Pen) ADG7TL3ENISF (D-Cys) - [NHZ] 64 Structure

ECSO(nM) ECSO(nM) (CSO(nM) Proliberanon Microphys.

Structure

TABLE 4

‡

(H)-(Pen)ADGPTLEEWISF (Cys)-(NH2)

[0-c-hh]-ldgtleemisf(cys)-(nh 2)

1 2 1

[H] - (Homocys) Addriftentsf [Homocys] - (NH 2]

ţ.

[O=C-NH]-\DGPTI\_RENISE (Homocys) - (NHZ]

ã

ţ

[H] - (Homocys) ADGFILHEWISF (Cys) - [NH2]

(0-C-H)-ADGPILRENISF-(Cys)-(NH<sub>2</sub>)

[0=C-NH]-ADGFTLAE#ISF(Pen)-(NH2)

ţ [O=C-NH]-EDGPTIREWISF(Cys)-[NH2] Ph-CE-

[H]-KADGPTLREWISFE-[NH2]

[H]-(Cys)ADGFTLREWISF(D-Cys)-[NH2]

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[H]-(D-Cys)ADGFTLREWISF(D-Cys)-[NHZ]

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In this example amino acid substitutes at positions D, E, I, S, or F in the cyclized compound

CADGPTLREWISFC

were assayed for  $\mathrm{EC}_{50}$  and  $\mathrm{IC}_{50}$  values as described above. Microphysiometer results are given in parentheses. The ទ

results are summarized in Table 5 below.

EXAMPLE 7

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(H)-FADGPILIENISER-(NH2)

0=C-NH-

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[0-C-NH]-ADGPILRENIEF(Cys)-[NH2]

ECSO(nM) ECES(nM) ICEO(nM) Proliferation Minagany.

Structure

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[H]-(Pen)ADGFTLREWISF(Pen)-[NH2]

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[HN]-ADGPILREMISFE-[NH2]

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**+** 

[O=C-NH]-ADGFTLRENISF(Cys)-[NH2]

‡

F: (DIPh - Ala) F - (1-Nal)

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TABLE 5

CADGPILREWISFC

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	IC50 (nM)	Q 2	Ŋ	<b>+</b>	‡	‡	‡	‡	‡	‡	÷	+	‡	ţ	<b>;</b>	‡	‡
	ECSO (n.M) Cell Projif.	a a	+	(+)·+	+	-1∙	<b>‡</b>	( <del>t</del> ) <del>t</del>	‡	1	+	( <del>1</del> ) ‡	1	( <del>‡</del> )‡	(÷) <b>‡</b> ‡	‡	( <del>‡</del> ) ‡
	Substitution	S - (Homoser)	S - (N-Me-Leu)	F-A	F - D-Ala	F - D.Phe	F - Homo-Phe	F. CHA	1.11	F - (Ser(Bzl))	F - (N-Me-Ala)	F • (Phenyigly)	F - (Pyridylala)	F - (p-Nitrophe)	F • (3,4-di-Cl-Phe)	F. (p-Cl-Phe)	F - (2-Nal)

Substitution	EC50 (nM) Cell Prolif.	ICSO (n.M.)	
Б-0	(+) ++	‡	
D-A	÷	‡	
I-A		. <b>†</b>	
∀-'s			_
:	<u>;</u>	Ι.	
3 - EAIA	+	1	
S - 5ar	+	ţ	
S-Aib	€ +	‡	
S - D-Ser	÷	ţ	
S - Nva	(±) ±	‡	
S-Abu	‡	‡	
S - (N-Me-Ala)	÷	÷	
S - (N-Me-Val)	+	1	
S - (N-Me-Ala) •	÷	. <b>÷</b>	
S - (Nor-Leu)	ţ	‡	
S - (t-Bu-Gly)	÷	‡	
S - (N-Ma-Sanita)		-	

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EXAMPLE 8

In this example, amino acid substitutions in the

compound

[O - C - NH] - A D G P T L R E W I S F (CYS)

were evaluated at positions D, S, or F as indicated in Table 6 below.  $EC_{50}$  and  $IC_{50}$  values were calculated as described above. Microphysiometer results are in parentheses.

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1050 (n.M.)

EC50 (nM) Cell Projif.

Substitution

0

F - (N-Me-Phe)

15

+

S.F - Ava (thioether) S.F. Ava (cys-cys) Q

÷+++++ £

ADG - deletion AD - deletion

Ava = H2N COOH

**†**· +

S.F. Ava

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TABLE 6

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EXAMPLE 9

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In this example  $EC_{50}$  and  $IC_{50}$  values were calculated as described above for the dimer compounds listed in Table 7 below. The cyclized monomer

CADGPTLREWISPC

is included as a comparison.

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[O=C-NH]-ADGPTLREWISF(Cys)

0.00

The compounds of Table 8 were inactive at the maximum concentration tested of 10µm.

In Table 9,  $\mathrm{EC}_{50}$  and  $\mathrm{IC}_{50}$  values determined as described above for cyclized and dimerized variants of I E G P I L R Q W L A A R A are compared. In Table 10, truncations of the dimer 15

ICSO (mM)

EC50 (nM) Cell Prolif.

Substitution

9

(F)

D-E

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(土) ‡

free acid form

C-term. Gly addition

(H) - I E G P T L R Q W L A A R A (Bala) K - (NH<sub>2</sub>) (H) - I E G P T L R Q W L A A R A

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described above. Microphysiometer results are given in are compared.  $\Sigma C_{50}$  and  $IC_{50}$  values were calculated as

parentheses.

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‡

S.F - Abu, DiPh-Ala

F - DIPh-Ala S-Abu

3

(±)± (<del>†</del> **‡** 

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PCT/US96/096		ICSD (nM)	‡	ţ.	‡	‡	ŧ	+	‡	4	4.	1	
		ECSO (nM) nvs. Protif.	† +	ţ	†·	ţ	‡	‡	† †	‡	4	÷	
74		ECSO Meropnys.	‡	2	‡	1	2	‡	2	‡	+	‡	
WO 96/40750			CAUGETIEST	[Ac] - Acprilentisec [Ac] - Acprilentisec	ADGPT TENTSEC	(Ac)-Igpilaemisec (Ac)-Igpilaemisec	[AC] -GFTLRENISFC     [AC] -GFTLRENISFC	<u> </u>	(Ac) - <u>Filedisco</u>     Ac) - <u>Filedisco</u>	PILREALISTC	(Ac)-TIRENISEC     (Ac)-TIRENISEC	TERENIEST   TERENIEST	
PCT/US96/09623			ECSO (nM) ICSO (nM) <u>ponys. <u>Prolif</u></u>	‡		; ‡ +		‡ ‡	‡ ‡	; ;		+	
WO 96/40750 73	TABLE 7		EC50 Meronics	0 	( HN) - DESTREYING - (HF) 0 0	++ (H) - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	(H)-iedpitromlaara (β-ala) k- (NH2)	(H)-CIEGETI RQMLAARA- (NH2)  (H)-CIEGETI RQMLAARA- (NH2)	(H)-GDGFILSENISF-(NH2)     GDGFILRENISF-(NH2)	(H) -5VQCGFTLRQWLAARWEL5-(NH)		(H) -+WGPTLRSGC- (NH2) (H) -+WGPTLRSGC- (NH2)	

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(H)-CIESPIE ROWLES-RAC- (NH2)

(H)-CINOCIAN-(H)

TABLE 8

[H]-CELINGLASEC-(NH2) (H)-CHSQLLAC-(H) [H]-CISIOMIAC-[NH2] [H]-CORADITMEC-(NH2)

(H)-FERRITAGMA-(H)

(H)-CINGFFORMANIC-(NH2)

(H)-CII.CONLONENC-(NH2)

(H)-CIETVWKLARNC-(NH<sub>2</sub>) (H)-CT.SEELAGOCC-(NH2)

[H]-CILREFIDETIRVC-[NH2] [H]-CGIZGPILSIMICC-(NH2)

(H)-CSIKEFIESGIANOC-(NH2) [H]-CTTVGPSINSWLTC-(NH2)

(H)-CILAEFLASGVECC-(NH2)

[H]-CILXEMLVSHEWNC-(NH2)

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TABLE 9

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TABLE 10

(H) -IESPIT #G#L335A (\$-314) K- (NH,) (H) - IEGFT SCHLAMBA

ICSO (n.M.) 9 ECSO (nM) Cell Prolif. + (Ac) - IEGFTLEGWLAAR-BA-KINE,) (Ac) - TEGFT SOWLANDA Sequence

IC50 (nM)

ECSO (nM)

-**|**-

‡.

N.D

(H)-SEGPTIAGNIASS-(H)

‡

**‡** 

N.D

(H)-CIEGPILEQWIASAC-(NB)

**‡** 

‡

‡.

[H]-INTERMINARA(B-ALA)K-[MH2]

(H)-IEGPILZOWLAARA

Š ‡ (H) - IEGPTINGULASA-BA-K (NH,) (H) - IEGZTL?GWLAAR ACINDESTRUBEL - (H)

4 9 **£** 8 (אני) -בקרבואמתבראונים- (אני) (H) - IEGPTINGADA-BA-K (KH.) (AC) -EGFTLEGWLAARA

2 **+** (H) -EGFILRQALSASA-BA-K (NRL,) (H) -EGFTT3QALASAA

**†** 

1

‡

(H)-CIEGPILROWLANZA-(NH2) (H)-CIEGETLEQWIELER-(MP)

(<del>1</del>) + (H)-EGPTIRONIARA-BA-KINN, (H) -EGFTL ROWLALR

2

‡ (אנאו א-גם-גינשמעדנים) - בפדעומעניים (Ac) - EGTT SOME XX

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(H) - EGFTLEGNELAN

+ (H) -EGPTISGALAA-BA-K(NH.)

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introduced at positions G, P, and W in the cyclized compound In this example various substitutions were

EXAMPLE 10

[H] - CADGPTLREWISFC - [NH2].

Table 11 lists examples of the substituted compounds that show TPO agonist activity. The substitutions abbreviated

2

## TABLE 11

in the table are as follows:

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NH, I	*	jev	i a	£	1	1 2	1	1		182	ď	Nal	Tr	ia N	P.T.	1	
[H]-CADGPTLREWISFC-{NH,	<b>a</b>	Нур(ОВп)	Hyp(OBn)	Pro	Pro	Hyp(OBn)	Pro	Pro	Hvp(OBn)	8	à		Pro	L-Tic	D-Tic	D-Tic	Hyp(OBn)
	9	Sar	Sar	Gly	Gly	Sar	Gaba	Cpr-Gly	Sar	Ģļ	'n		iBo	Cpr-Gly	Gly	Cpr-Gly	Gabs

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Proline Replacements

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1-4-Hvo (OBn)

L-Azetidine carboxylic acid

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Tryptophan Replacements

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F.W. COO.

N COOH

B-alanine

COOH NA,

C.2.Nal

осу — 1 - Осу —

HAN COOH

BL-1-Mc-Trp

BL-3-Br-Trp

Mut,

HIN NHY DLS-F-Trp

Amine DL-5-Me-Trp

S NH2

L-(Berzonticny))-alamine

N-Pentyl glycine

N-Cyclopropyl glycine

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## EXAMPLE 11

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receptor, have been done. First, marrow cells, harvested from non-adherent cells that stain for AChE. These cells appear to incubation period, the cultures were concentrated by Cytospin, megakaryocytes), and counted by microscopic analysis. One (1) designated as "maximal". Control cultures containing no added growth factors produced very few AChE-positive cells. Several This finding was the first evidence that this peptide solid medium (methylcellulose) containing either no factors, 1 the femurs of 8 to 9 week one Balb/C mice, were incubated for total marrow cells/ml (in 50 ml cultures) an estimated 1 to 2 stained for acetylcholinesterase (AChE, a diagnostic of mouse of the peptide compounds were tested at high concentration in experiment, marrow cells were harvested and cultured in semicounted and grouped into small colonies (3-5 cells) or large negative control cultures. This indicates that the peptides To assess the feasibility of mice as a convenient mimic TPO in their ability to stimulate the expansion of the nM rhuTPO gave rise to the outgrowth of very large (>40 um) colonies of large cell (presumed to be megakaryocytes) were Peptide A at 10uM produced a maximal response of the mouse colonies (greater than 6 cells). The results are shown in be mature megakaryocytes. From an initial seeding of  $10^6\,$ imes 10 $^6$  megakaryocytes developed. This responce to TPO was test species, several in vitro experiments, designed to measure the activity of the test compounds on the mouse concentrations of the test peptides. At the end of the 7 days in liquid culture with either rhuTPO or various nM rhuTPO, or 10uM Peptide A. After 7 days in culture, this assay and the results are summarized in Table 12. substantially more colonies of both sized than did the family is active on the murine receptor. In a second TPO and the test peptides both produced Mk precursor cell population. Table 13. marrow.

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activity of the test compounds on murine and human receptors, To obtain a more quantitative comparison of the

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the muTPO receptor was cloned and transfected into BaF3 cells.

A TPO dependent population of cells was isolated.

## TABLE 12

s

Peptide	Concentration mosts	
Ω	100 000	Ind) Regponse
	200,100=	none
O.	40,000	maxima]++
C + S.A. *	1000	meximal at
		- Tellityou
S.A. alone	1000	none
В	100.000	
		Thurst
4	10,000	maximal**
TPO (R & D)		
		"maxima]"

2

 Streptavidin complexed to biotinylated peptide - concentration of putative 1:4 complex.

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\*\* Compared to recombinant human TPO

\*\* 25-30% ACE staining cells on cytopspin

No factor cultures - ca. 5% AChE staining cells (lower cellularity)

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## TABLE 13

Compound         3-5 large compound           No factors         1         2           No factors         2         1           1 nM TPO         #1-1         15           1 nM TPO         #2-1         12           1 nM TPO         #2-1         16           1 nM TPO         #2-2         13           10 uM Peptide         #1-1         25           10 uM Peptide         #2-1         25           10 uM Peptide         #2-1         22	ells 6-12 large cells	0					2 6	7 5	Q.	0 6	
Pun	3-5 large cells	2	1	15	12	16	13	25	22	22	
	Compound	No factors 1	No factors 2								

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The disclosures in this application of all articles and references, including patent documents, are incorporated herein by reference in their entirety for all purposes.

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IN THE CLAIMS

receptor	
thrombopoietin	
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binds	
that	
punoduo	having:
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1.	compor
	said

(1) a molecular weight of less than about 8000

daltons, and

(2) a binding affinity to thrombopoietin receptor as expressed by an  ${\rm IC}_{50}$  of no more than about 100  $\mu{\rm m}$ . The compound of Claim 1, wherein said compound is a

peptide, and,

wherein from zero to all of the -C(O)NH- linkages of the peptide have been replaced by a linkage selected from the group consisting of a

-CH2OC(O)NR- linkage; a phosphonate linkage; a -CH2S(O)2NRlinkage; a -CH<sub>2</sub>NR- linkage; and a -C(0)NR<sup>6</sup>- linkage; and a

-NHC(O)NH- linkage where R is hydrogen or lower alkyl and  $\mathrm{R}^6$ is lower alkyl, further wherein the N-terminus of said peptide or peptide group; a -NRC(0)R group; a -NRC(0)OR group; a -NRS(0)2R group; mimetic is selected from the group consisting of a  $-NRR^{\perp}$ 

benzyloxycarbonyl-NH- group; and a benzyloxycarbonyl-NH- group having from 1 to 3 substituents on the phenyl ring selected a -NHC(0)NHR group; a succinimide group; a 13

chloro, and bromo, where R and R1 are independently selected from the group consisting of lower alkyl, lower alkoxy, from the group consisting of hydrogen and lower alkyl, 18

and -NR<sup>3</sup>R<sup>4</sup> where R<sup>3</sup> and R<sup>4</sup> are independently selected from the and still further wherein the C-terminus of said peptide group of the N-terminus of the peptide so as to form a cyclic nitrogen atom of the  $-NR^{\rm J}R^4$  group can optionally be the amine selected from the group consisting of hydroxy, lower alkoxy, group consisting of hydrogen and lower alkyl and where the or peptide mimetic has the formula -C(0)R2 where R2 is

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and physiologically acceptable salts thereof.

peptide,

 A pharmaceutical composition comprising the compound of Claim 1 in combination with a pharmaceutically acceptable carrier.

4. A method for treating a patient suffering from a disorder that is susceptible to treatment with a thrombopoietin agonist, comprising admistering to the patient, a therapeutically effective dose or amount of a compound of Claim 1.

administered to the patient is a peptide, and,
wherein from zero to all of the -C(O)NH- linkages of the

peptide have been replaced by a linkage selected from the group consisting of a -CH<sub>2</sub>OC(0) NR- linkage; a phosphonate linkage; a -CH<sub>2</sub>NR- linkage; and a -C(0) NR<sup>6</sup>- linkage; and a -NHC(0) NH- linkage where R is hydrogen or lower alkyl and R<sup>6</sup> is lower alkyl,

further wherein the N-terminus of said peptide or peptide mimetic is selected from the group consisting of a -NRR1 group; a -NRC(0)R group; a benzyloxycarbonyl-NH- group; and a benzyloxycarbonyl-NH- group; and a benzyloxycarbonyl-NH- group having from 1 to 3 substituents on the phenyl ring selected from the group consisting of lower alkyl, lower alkoxy, chloro, and bromo, where R and R<sup>1</sup> are independently selected from the group consisting of hydrogen and lower alkyl,

and still further wherein the C-terminus of said peptide or peptide mimetic has the formula -C(0)R<sup>2</sup> where R<sup>2</sup> is selected from the group consisting of hydroxy, lower alkoxy, and -NR<sup>3</sup>R<sup>4</sup> where R<sup>3</sup> and R<sup>4</sup> are independently selected from the group consisting of hydrogen and lower alkyl and where the nitrogen atom of the -NR<sup>3</sup>R<sup>4</sup> group can optionally be the amine group of the N-terminus of the peptide so as to form a cyclic peptide,

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and physiologically acceptable salts thereof.

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7. The compound of claim 6, wherein said sequence of amino acids is cyclized. 8. The compound of claim 6, wherein said sequence of

amino acids is dimerized.

9. The compound of Claim 6, wherein the compound comprises the sequence of amino acids

comprises the sequence of amino acids  $C X_2 X_3 X_4 X_5 X_6 X_7$  where  $X_2$  is K, L, N, Q, R, S, T or V;  $X_3$  is C, F, I, L, M, R, S or V;  $X_4$  is any of the 20 genetically coded L-amino acids;  $X_5$  is A, D, E, G, S, V or Y;  $X_6$  is C, F, G, L, M, S, V, W or Y; and  $X_7$  is C, G, I, K, L, M, N, R or V.

10. The compound of Claim 8, wherein  $X_4$  is A, E, G, H, K, L, M, P, Q, R, S, T, or W.

11. The compound of Claim 10, wherein  $X_2$  is S or I;  $X_3$  is L or R;  $X_6$  is R;  $X_5$  is D, E, or G;  $X_6$  is F, L, or W; and  $X_7$  is I, K, L, R, or V.

12. The compound of Claim 9, wherein said compound comprises a sequence of amino acids:  $X_8 \subset X_2 \times_3 X_4 \times_5 X_7$ 

where  $X_2$  is F, K, L, N, Q, R, S, T or V;  $X_3$  is C, F, I, L, M, R, S, V or W;  $X_4$  is any of the 20 genetically coded L-amino acids;  $X_5$  is A, D, E, G, K, M, Q, R, S, T, V or Y;  $X_6$  is C, F, G, L, M, S, V, W or Y;  $X_7$  is C, G, I, K, L, M, N, R or V; and  $X_8$  is any of the 20 genetically coded L-amino acids.

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ò The compound of Claim 12, wherein  $X_{\theta}$  is G, S, Y . 13

GGCTLREWLHGG The compound of Claim 12, wherein said compound comprises a sequence of amino acids: 14. FCGG.

The compound of Claim 6, wherein said compound comprises a sequence of amino acids: 5.

X<sub>8</sub> G X<sub>1</sub> X<sub>2</sub> X<sub>3</sub> X<sub>4</sub> X<sub>5</sub> W X<sub>7</sub>

V, or W; X, is A, K, L, M, R, S, V, or T; X5 is A, E, G, K, M, where X<sub>1</sub> is L, M, P, Q, or V; X<sub>2</sub> is F, R, S, or T; X<sub>3</sub> is F, L, Q, R, S, or T; X, is C, I, K, L, M or V; and X<sub>8</sub> is any of the

20 genetically coded L-amino acids.

The compound of Claim 15, wherein  $x_1$  is P;  $x_2$  is T; X<sub>3</sub> is L; X<sub>4</sub> is R; X<sub>5</sub> is E or Q; X<sub>7</sub> is I or L. 16.

4 4

The compound of Claim 16, wherein said compound 17.

comprises a sequence of amino acids:

X9 X8 G X1 X2 X3 X4 X5 W X7

where X<sub>8</sub> is A, C, D, E, K, L, Q, R, S, T, or V; and X<sub>9</sub> is A, C, E, G, I, L, M, P, R, Q, S, T, or V.

ĸ 占 m The compound of Claim 17, wherein  $X_{\theta}$  is D, and X<sub>9</sub> is A or I. 18.

selected from the group consisting of GGCADGPTLREW GPTLREWISFCGGK; TIKGPTLRQWLKSREHT S; SIE GPTLREWLTSRTPHS; LAIEGPTLROW I S F C G G; G N A D G P T L R Q W L E G R R P K N; G G C A D LHGNGRDT; CA.DGPTLREWISFC; and IEGPTL The compound of Claim 18, wherein said compound is OWLAARA. 19.

The method of Claim 4, wherein said compound that is administered to the patient comprises a sequence of amino 20. acids:

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C X<sub>2</sub> X<sub>3</sub> X<sub>4</sub> X<sub>5</sub> X<sub>6</sub> X<sub>7</sub>

where X<sub>2</sub> is K, L, N, Q, R, S, T or V; X<sub>3</sub> is C, F, I, L, M, R, S or  $V_i$   $X_4$  is any of the 20 genetically coded L-amino acids;

Xs is A, D, E, G, S, V or Y; Xg is C, F, G, L, M, S, V, W or

Y; and X, is C, G, I, K, L, M, N, R or V.

Н, К, ຶ່ 21. The method of Claim 20, wherein  $X_4$  is A, E,

M, P, Q, R, S, T OT W. ij

22. The method of Claim 21, wherein  $X_2$  is S or T;  $X_3$ 

Lor R; X<sub>4</sub> is R; X<sub>5</sub> is D, E, or G; X<sub>6</sub> is F, L, or W; and X<sub>7</sub> is I, K, L, R, or V.

23. The method of Claim 22, wherein said compound that

is administered to the patient comprises a sequence of amino acids: GGCTLREWLHGGFCGG.

24. The method of Claim 4, wherein the disorder

susceptible to treatment with a thrombopoietin agonist is

selected from the group consisting of:

chemotherapy, radiation therapy, or bone marrow transfusions hematological disorders and thrombocytopenia resulting from

The method of Claim 4, wherein said compound that is 25.

administered to the patient comprises a sequence of amino

acids:

X<sub>8</sub> G X<sub>1</sub> X<sub>2</sub> X<sub>3</sub> X<sub>4</sub> X<sub>5</sub> W X<sub>7</sub>

where X<sub>1</sub> is L, M, P, Q, or V; X<sub>2</sub> is F, R, S, or T; X<sub>3</sub> is F,

V, or W; X, is A, K, L, M, R, S, V, or T; X5 is A, E, G, K, M,

Q, R, S, or T; X, is C, I, K, L, M or V; and X<sub>8</sub> residue is any

of the 20 genetically coded L-amino acids.

26. The method of Claim 25, wherein  $X_1$  is P;  $X_2$  is T;  $X_3$ 

is L; X4 is R; X5 is E or Q; X7 is I or L.

27. The method of Claim 26, wherein said compound comprises a sequence of amino acida:

X<sub>9</sub> X<sub>6</sub> G X<sub>1</sub> X<sub>2</sub> X<sub>3</sub> X<sub>4</sub> X<sub>5</sub> W X<sub>7</sub>

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where Xg is A, C, D, E, K, L, Q, R, S, T, or V; and Xg is A, C, E, G, I, L, M, P, R, Q, S, T, or V.

X, The method of Claim 27, wherein  $X_8$  is D, E, or and X9 is A or I. 28.

PTLRQWLEGRRPKN; GGCADGPTLREWISFCG The method of Claim 28, wherein the compound that is consisting of GGCADGPTLREWISFCGG; GNADG G K; TIK G P T L R Q W L K S R E H T S; S I E G P T L R E W LTSRTPHS; LAIEGPTLROWLHGNGRDT; CAD GPTLREWISFC; and IEGPTLROWLAARA. administered to the patient is selected from the group

212

wherein said compound is selected from the group consisting of 30. A compound that binds to thrombopoietin receptor,

CADGPTLREWISFC

- [amide] U £4 - CADGPTLREWIS [Ac]

٠. NH<sub>2</sub> O = CADGPTLREWISFC

111 112 113 114 115 115

and

IEGPTLRQWLAARA (Bala)-K (NH<sub>2</sub>) IEGPTLROWLAARA

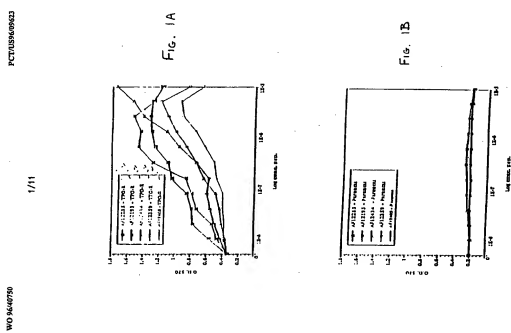
pateint a compound is selected from the group consisting of 31. A method for treating a patient suffering from a thrombopoietin agonist, comrpising administering to the disorder that is susceptible to treatment with a

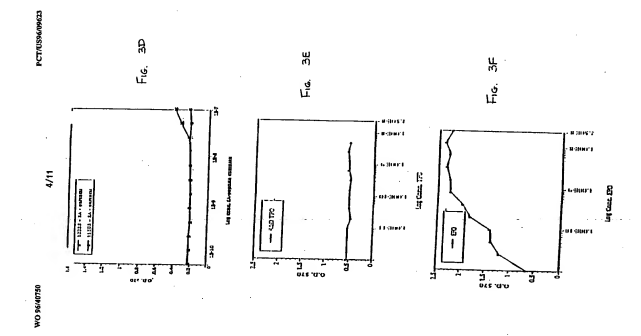
CADGPTLREWISFÇ

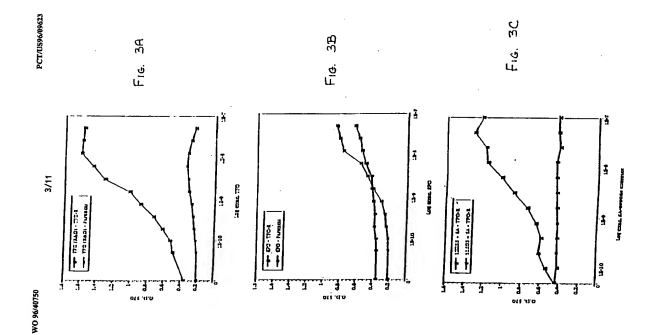
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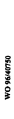
PCT/US96/09623 IEGPTLROWLAARA (Bala)-K [NH2] IEGPTLROWLAARA [Ac] - CADGPTLREWISFC - [amide] O = CADGPTLREWISFC - NH2 92 H<sub>2</sub>

and









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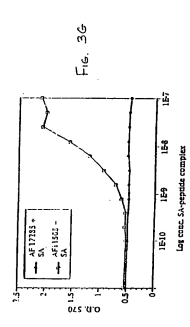
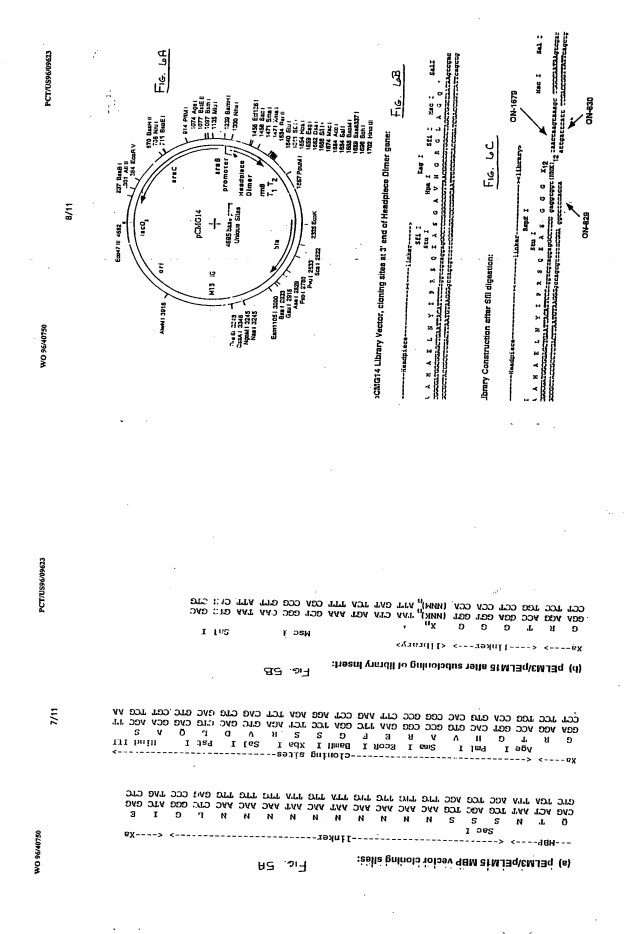


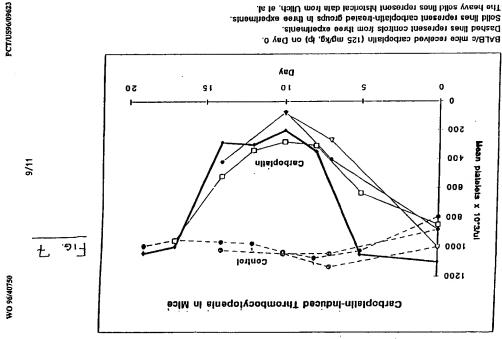
Fig. 4A 1684 AOR 1 1884 ELD12D? 1074 Age 1 : 691 Bd. 214 PIDM ! 2286 Ears)
2216 Ears)
2216 Ears)
2217 Ears)
2218 Ears) 736 Nau 1 Test base pairs Unto.:: Siles 1622 1 SDN Fe see Earn 1051 13011 

Fig. 4B pJS142 Library Vector, cloning sites at 3' end of lact gene:

Kec Library Offgo library Consunction after SIII digestion:

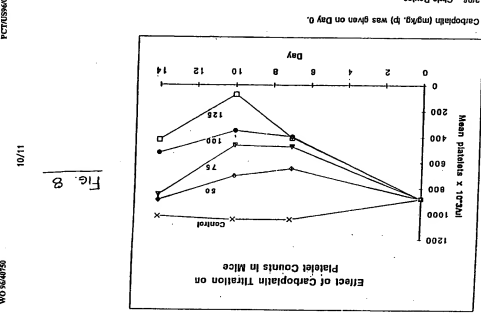
Fig. 4C





BALB/c mice received carbopialin (125 mg/kg, ip) on Day 0. Solid lines represent controls from three experiments. Solid lines represent carbopiatin-treated groups in three experiments. The heavy solid lines represent historical data from Ulich, et al.

3/96 Chris Boytos



3/96 Chits Boylos

PCT/US96/09623 11/11 F16. WO 96/40750

513+CBP 50 513+CBP 513+CBP 100 513+CBP 125 CBP 100 CBP 125 엹 g Control 513 5 3 75 Mean platelets x 10"3/ul 001 009 008 0001 1500 1400 0091 Tiltailon of Carboplatin Thrombocytopenia on Day 10 by AF12513: Amelloration of Carboplatin-induced

3/96 Chils Boylos Carboplalin (CBP; 125-50 mg/kg, ip) was given on Days 1-9. AF12513 (513; 196 $^{\circ}$  mg/kg, ip) was given on Days 1-9.

INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/09623

. CLASSIFICATION OF SUBJECT MATTER PC(6) :COTK 702, 706, 706, 7159, 7154, 1452 US CL. : 514/09, 11, 13, 14, 15, 16; 530/317, 323, 326, 327, 328, 329 Weredring to International Patent Charification (IPC) or to both militari charification and IPC FIELDS SEARCHED

inimum documentation searched (classification system followed by classification symbols)

U.S. : \$14/09, 11, 13, 14, 15, 16; \$30/317, 323, 326, 327, 328, 329

umentation scarched other than minimum documentation to the extent that such documents are included in the fields scarched

ilectronic data base consulted during the international search (name of data base and, where practicable, search terms used) APS, DIALOG, MEDLINE search terms: thrombopoletin, ligand, fregment, agonist, antagonist, receptor, mimetic, inhibit

DOCUMENTS CONSIDERED TO BE RELEVANT

Relevant to claim No. 1:31 US 5,358,934 A (BOROVSKY ET AL) 25 October 1994 (25.10.94), see column 2, line 64-column 3, line 8. US 5.141,851 A (BROWN ET AL.) 25 August 1992 (25.08.92), see column 9, lines 40-52. US 5,411,942 A (WIDMER ET AL.) 02 May 1995 (02.05.95), see column 5, line 37-column 6, line 11. Citation of document, with indication, where appropriate, of the relevant passages Category

<u>ن</u> WO 96/17062 A1 (ZYMOGENETICS,INC.) 06 June 1996 (106.06.96), see entire document.

See patent family annea Further documents are listed in the continuation of Box C.

Date of mailing of the international search report 1 7 SEP 1998 filling date but hier than Date of the actual completion of the international search 23 AUGUST 1996

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INTERNATIONAL SEARCH REPORT

Relevant to claim No. International application No. PCT/US96/09623 1:3 1-3 KATO et al. Purification and Characterization of Thrombopoietin.

J. Biochem. 1995, Vol. 118, pages 229-236, see entire document. WADA et al. Characterization of the Truncated Thrombopoietin Variants. Biochemical and Biophysical Research Communications. 24 August 1995, Vol. 213, No. 3, pages 1091-1098, see entire document. C. (Continuation), DOCUMENTS CONSIDERED TO BE RELEVANT
Cacgogy\*

Citation of document, with indication, where appropriate, of the relevant passages Α, Ρ

Form PCT/ISA/210 (continuation of account sheet)(July 1992)\*

## PATENTS · TRADEMARKS · DESIGNS · COPYRIGHT

## **European Patent Register Extract**

EUROPEAN PATENT REGISTER / EPIDOS

24/09/2002

-485301

AN:

96919241.8 PN: 0885242 IPC: C07K7/02

PART I - REGISTER OF EUROPEAN PATENTS (R92 EPC)

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APPLICATION NUMBER

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SUPPLEMENTARY SEARCH REPORT

INTERNATIONAL APPLICATION NR. INTERNAT. PUBL. NUMBER AND DATE

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PRIORITY

CLASSIFICATION

DESIGNATION

TITLE

APPLICANT

INVENTOR

: 0885242

: 96919241.8

: 19.12.1996/A1 : 19.12.1996

: 04.02.2000 : US9609623

: WO96040750 19.12.1996

: 07.06.1996

: 07.06.1995/ US 07.06.1995/ US

478128 : C07K7/02, C07K7/06, C07K7/08, C07K7/50,

C07K7/54, C07K14/52

: AT BE CH DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

: PEPTIDES AND COMPOUNDS THAT BIND TO A

THROMBOPOIETIN RECEPTOR

: FOR : ALL DESIGNATED STATES

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: FILLER, WENDY ANNE, DR., ET AL

GLAXOSMITHKLINE CORPORATE INTELLECTUAL PROPERTY (CN9.25.1) 980 GREAT WEST ROAD

BRENTFORD, MIDDLESEX TW8 9GS/GB

REQUEST FOR EXAMINATION : 22.12.1997

PART II - INFORMATION REGISTER (EPIDOS)

THIS APPLICATION IS BEING TREATED IN (/FAX-NR): MUNICH

/(+49-89) 23994465

THE PROCEDURE LANGUAGE IS (DE/EN/FR) : EN

PCT - CHAPTER II

REPRESENTATIVE

REQUEST FOR PRELIMINARY EXAMINATION

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: 12.11.1997

ACTS PERFORMED FOR ENTRY INTO THE REGIONAL PHASE

- NATIONAL BASIC FEE PAID

: 12.11.1997

died Marie Bride Strike

## ERIC POTTER CLARKSON

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## **European Patent Register Extract**

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- SEARCH FEE PAID
                                        : 31.12.1997
- DESIGNATION FEE(S) PAID
                                        : 12.11.1997
- EXAMINATION FEE PAID
                                       : 22.12.1997
CHAPTER - EXTENSION OF THE PATENT
COUNTRY-CODE / DATE PAYMENT / DATE WITHDRAWAL : AL/12.11.1997/00.00.0000
COUNTRY-CODE / DATE PAYMENT / DATE WITHDRAWAL : LT/12.11.1997/00.00.0000
COUNTRY-CODE / DATE PAYMENT / DATE WITHDRAWAL : LV/12.11.1997/00.00.0000
COUNTRY-CODE / DATE PAYMENT / DATE WITHDRAWAL : SI/12.11.1997/00.00.0000
 CHAPTER - RENEWAL FEES (ART.86)
 RENEWAL FEE A.86 (PATENT YEAR/PAID)
                                       : 03/15.06.1998
                                         04/14.06.1999
                                         05/13.06.2000
                                         06/13.06.2001
                                         07/12.06.2002
CHAPTER - CITED DOCUMENTS
THIS CHAPTER SHOWS THE ACTUAL SITUATION OF THE CITED DOCUMENTS.
NO OBLIGATION IS TAKEN FOR THE COMPLETENESS OF ALL THE CASES.
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PA:WO A 9 521 919
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A : J. BIOCHEM., 1995, VOL. 118, KATO ET AL., "PURIFICATION AND
   CHARACTERIZATION OF THROMBOPOIETIN", PAGES 229-236.
AP :BIOCHEMICAL AND BIOPHYSICAL RESEARCH COMMUNICATIONS, 24 AUGUST 1995, VOL.
   213, NO. 3, WADA ET AL., "CHARACTERIZATION OF THE TRUNCATED THROMBOPOIETIN
   VARIANTS", PAGES 1091-1098.
** END OF DATA **
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